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(54) Title: SYNTHETIC POLYNUCLEOTIDES

(57) Abstract

This invention relates to synthetic polynucleotides which encode lysyl oxidase, lysyl oxidase like molecules or variants of these species. The synthetic polynucleotides of the invention permit the expression of functional lysyl oxidase, lysyl oxidase like molecules or variants of these species, typically in good yield. The invention also relates to recombinant DNA molecules containing these, synthetic polynucleotides, to cells containing them and to uses of the expression products.

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"Synthetic Polynucleotides"

TECHNICAL FIELD

The present invention relates to synthetic polynucleotides encoding proteins and polypeptides selected from the group consisting of lysyl oxidases, lysyl oxidase-like proteins and variants thereof; to use of the synthetic polynucleotides to provide high yields of those proteins and polypeptides; and to uses of the proteins and polypeptides produced from the synthetic polynucleotides.

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BACKGROUND ART

Lysyl oxidase is a copper dependent amine oxidase that catalyses the oxidation of amines, including but not limited to primary amines and, in particular, the amine side chain of lysine. Lysine oxidation catalysed by lysyl oxidase has been observed in the oligopeptide and polypeptide chains of collagen and tropoelastin. Lysyl oxidase activity has been observed with other amine containing substrates, such as oligopeptides where the efficiency of catalyzed oxidation is dependent on adjacent sequences (Kagan et al., 1995a) such as vicinal dicarboxylic amino acid residues (Nagan and Kagan, 1994).

With varying efficiencies, lysyl oxidases can oxidise other substrates, such as butylamine and phydroxybenzylamine, for example, to form butyraldehyde and phydroxybenzylaldehyde respectively. Non-peptide reactivity is also displayed with, for example, semicarbazide(s) and oxidation of tyramine (Palcic et al., 1995). Lysyl oxidase activity has also been observed with other amine containing molecules including inhibitors and unnatural substrates such as trans-2-phenylcyclopropylamine (Shah et al., 1993).

The majority of the work done to date in characterising lysyl oxidase has been with respect to non-human mammalian lysyl oxidases. The level of amino

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acid homology between lysyl oxidases from different species is of the order of 90%.

"Lysyl oxidase-like proteins" have also been identified by the analysis of nucleotide and predicted amino acid sequence alignments of DNA and protein molecules which are expressed in a particular mammalian species including humans. The level of homology between these proteins and lysyl oxidases is of the order of 75%, which is a highly significant degree of homology. Evidence indicates that these molecules may function as

enzymes in the extracelluar space as members of a lysyl oxidase family of molecules, and in the cell to provide lysyl oxidase activities (Kim, Y. et al. 1995).

The lysyl oxidase gene encodes a single polypeptide species and there has been no observation of lysyl oxidase mRNA splice variants or isoforms of the lysyl oxidase polypeptide in vivo (Boyd et al., 1995).

In the production of intra- and inter-molecular crosslinked molecules, lysine oxidation in tropoelastin is a necessary step in the formation of allysine, desmosine and isodesmosine condensation products.

Crosslinked molecules, including elastin and collagen, are significant components of fully functional connective tissue. In this regard, deficiencies in lysyl oxidase such as that found in lathyrism, lead to marked phenotypic changes that can compromise the viability of an individual.

Lysyl oxidase exists in at least two forms: an intracellular protein and the more thoroughly characterised extracellular form. In the intracellular form its roles include a ras recission function, and the encoding gene has been classified in this context as a ras-recission gene or rrg (Contente et al., 1990; Kenyon et al., 1991). In its capacity as rrg, its expression is altered in an incompletely catalogued manner, to apparently reduce the oncogenic phenotype of cells expressing aberrant ras. Lysyl oxidase levels also change during differentiation and development

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(Dimaculangan et al., 1994) and in response to growth factors (Green et al, 1995). Lysyl oxidase is also a secreted protein, available in the extracellular matrix of some connective tissues in very low concentrations.

Difficulties in obtaining sufficient quantities of enzyme for biochemical analysis have impeded detailed exploration of its properties. Yields of purified naturally occurring lysyl oxidase available from typical purification procedures have been limited to 2-4mg starting with 0.5 to 1kg of cleaned bovine aortae (Kagan and Cai, 1995).

Extracellular lysyl oxidase is typically made as a larger protein, which includes a collection of amino acid residues at the amino terminus of the protein. This form, termed preprolysyl oxidase is secreted—from the cell in a process contemporaneous with cleavage of the amino-terminal region to generate prolysyl oxidase which in turn is cleaved outside the cell to generate the mature form of the protein.

20 The extracellular (and possibly intracellular) lysyl oxidase enzyme additionally contains copper and at least one organic cofactor which is postulated to be a quinonelike component. The organic cofactor has variously been considered to be covalently or non-covalently bound 25 to the enzyme (Williamson et al., 1986; Kagan and Trackman, 1991). Examples of organic cofactors, which have been found to be associated with oxidases, include quinones such as P.Q.Q., topa quinone (2,4,5 trihydroxyphenylalanine quinone) and tryptophan 30 tryptophylquinone (Tanizawa, 1995). The organic cofactor for lysyl oxidase is now known to be a quinone. possible that the cofactor can be supplied, generated by interaction of a quinone derivative with the depleted apo-form of the enzyme, and/or generated by protein 35 oxidation such as that mediated by the participation of copper. There is at least one atom of copper tightly bound per one molecule of functional enzyme. However it was experimentally demonstrated (Gacheru et al., 1990)

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that when copper is removed by a chelating agent the inactive apoenzyme can be restored to its former level of activity by the addition of copper.

Posttranslational modification(s) such as glycosylation have not been described for the mature form of lysyl oxidase, although there is evidence for glycosylation of the amino terminal region destined for removal during maturation of the precursor prolysyl oxidase (Cronshaw et al., 1995). As other forms of modification exist in vertebrate cells, and include participation by the cytosol, Golgi and secretion machinery, followed by extracellular processing, uncertainty has surrounded the question of whether a functional lysyl oxidase could be made in host cells other than verterbrate systems.

Isolation of endogenous lysyl oxidase from, for example, mammalian tissue typically uses chemical agents which interfere with protein association, such as urea. Modest amounts of lysyl oxidase can be recovered in this way from skin (Shackleton and Hulmes, 1990). The purified material is typically maintained in solutions containing chemical agent(s), in part to minimise protein aggregation and loss of catalytic function. When prepared in this way, the enzyme displays sluggish activity, which has led to the assertion that the enzyme (or a macromolecular complex) (Cronshaw et al.,1993) is altered during the relatively harsh extraction procedure (Shackleton and Hulmes, 1990).

To circumvent difficulties associated with the use of the purified naturally occurring enzyme, recombinant production is a logical alternative. However despite the fact that inferred lysyl oxidase sequences including human (Mariani et al., 1992; Svinarich et al., 1992; Kenyon et al., 1993), rat (Trackman et al., 1990) and mouse (Contente et al., 1993) have been available for several years there have been no reports in the literature of production of recombinant functional lysyl oxidase using non-mammalian hosts for other than modest

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levels of expression and no functional expression of the human sequence. Kagan et al. (1995b), have reported the production of modest levels of porcine lysyl oxidase from cDNA in mammalian cells indicating that bacterial production of functional enzyme was problematic. One of the impediments to the production of recombinant lysyl oxidase has been the discrepancies appearing in some of the available sequence information (e.g. Trackman et al., 1990; Hamalainen et al., 1991 vs Mariani et al., 1992).

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10 DESCRIPTION OF THE INVENTION

The paucity of information in the literature in relation to the nature of host-specific post-translational modifications of lysyl oxidase (e.g. Cronshaw et al. 1995), together with the real absence of reports of functional lysyl oxidase production in non-mammalian recombinant expression systems, teaches against the choice of non-mammalian hosts for expression of nucleotide sequences encoding mammalian lysyl oxidase and implies that non-mammalian host cells are unlikely to produce functional lysyl oxidase which is encoded by a mammalian nucleotide sequence.

The present inventor has reasoned that expression of mammalian lysyl oxidase in non-mammalian host cells might be achieved using a synthetic polynucleotide which encodes lysyl oxidase but which has been specifically designed for use in non-mammalian host cells such as (but not limited to) Escherichia coli, Saccharomyces cerevisiae and Pichia. In particular, the inventor proposed the hypothesis that non-mammalian host cells have difficulty expressing from nucleotide sequences encoding mammalian lysyl oxidase because of the presence of 'rare' codons in those sequences which are used infrequently in the non-mammalian host cell. Rare codons are codons that are found at low frequency in highly expressed genes in a particular host. A measure of efficiencies of codon usage is the codon adaptation index (CAI) (Sharp and Li 1987). Highly expressed genes such

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as those encoding ribosomal proteins in *E. coli* typically have high CAI values that are greater than or equal to 0.45, whilst poorly expressed genes have much smaller CAI values; for example, the *E. coli lacI* gene has a CAI of 0.3

The inventor calculated the CAI of a nucleotide sequence encoding a secreted form of human lysyl oxidase for a number of non-mammalian hosts by considering the usage of codons in the nucleotide sequence shown in Figure 1. The CAI is calculated to be 0.24 for E. coli and 0.11 for Saccharomyces cerevisiae. The inventor identified codons in the nucleotide sequence encoding human lysyl oxidase which are infrequently used in host cells such as E. coli including: an AGGAGA cluster encoding adjacent arginine residues at positions 937 to 942 which has the capacity to limit expression and generate translational pausing and frameshifting in E. coli; 11 other instances of single rare AGG and AGA arginine codons capable of instigating translational hopping (Kane 1992); and an ensemble of codons rarely found in highly expressed E. coli genes (Table 1). Together these codons account for 28% of the amino acid coding capacity of the sequence.

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Codons found in the natural lysyl oxidase sequence of Figure 1 which are rarely found in highly expressed E. coli genes

5	Codon	Amino acid	Frequency in highly expressed genes %	Occurrence 1 residues (% total se		
	AGG	Arg	0.14	6	2.4	
	AGA	Arg	0.21	7	2.8	
	CGA	Arg	0.31	2	0.8	
	ccc	Pro	0.43	7	2.8	
.0	TGT	Cys	0.47	7	2.8 _	
	TGC	Cys	0.61	3	1.2	
	ACA	Thr	0.65	7	2.8	
	CCT	Pro	0.66	6	2.4	
	TCA	Ser	0.68	3	1.2	
.5	GGA	Gly	0.70	4	1.6	
	AGT	Ser	0.72	6	2.4	
	CCA	Pro	0.82	2	0.8	
	TCC	Ser	0.94	4	1.2	
	GGG	Gly	0.97	2	0.8	
0	CTG	Leu	0.99	6	2.4	

In consideration of the CAI values for the nucleotide sequence encoding human lysyl oxidase in E. coli and Saccharomyces cerevisiae, the inventor reasoned that the problem associated with expression of verterbrate lysyl oxidase or lysyl oxidase-like molecules in particular host cells or expression systems was directly related to the high proportion of codons in the verterbrate lysyl oxidase or lysyl oxidase-like protein nucleotide sequence which are rarely used by the particular host cells. The inventor further reasoned that verterbrate lysyl oxidase and lysyl oxidase-like molecules might be expressed in alternative host cells if

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the nucleotide sequences encoding those molecules were able to be specifically designed so as to favour higher levels of expression of lysyl oxidase in a particular alternative host cell. The inventor reasoned that nucleotide sequences which encode verterbrate lysyl oxidase or lysyl oxidase-like proteins and which have been specifically designed for high level expression in a particular host may also be suitable for expression in other host cells provided that the specifically designed nucleotide sequence contained codons which were frequently used by the other host cells. Moreover, the inventor realised that the concept of modifying the nucleotide sequence would find application in producing high level expression of lysyl oxidase or lysyl oxidaselike proteins in the natural host cell, in particular, high level expression of nucleotide sequences encoding human lysyl oxidase and lysyl oxidase-like proteins in human cells.

The present invention provides high level expression of nucleotide sequences encoding lysyl oxidases and lysyl oxidase-like proteins in recombinant expression systems. The inventor has recognised that lysyl oxidases, lysyl oxidase-like proteins and variants thereof can be used in a variety of, for instance, pharmaceutical applications, but these uses require significant quantities of lysyl oxidase, lysyl oxidase-like proteins or variants thereof in a native conformation. Thus the realisation of the potential application of lysyl oxidase, lysyl oxidase-like expression products and variants thereof first requires a system which allows high level production of those products in native conformation.

The benefits of being able to produce significant amounts of expression product in a functional conformation include specific oxidation, rather than random oxidation, so as to produce cross-linked complexes including cross-linked elastin or cross-linked collagen which are structurally and functionally similar to normal physiological complexes. Secondly, the expression

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product which exists in a functional or native conformation is unlikely to cause an adverse reaction in vivo, whether the expression product is directly administered in vivo or in the situation where residual expression product remains in a cross-linked complex after catalysis, wherein that complex is intended for in vivo use.

The invention provides for high level expression of vertebrate nucleotide sequences encoding lysyl oxidase and lysyl oxidase-like proteins by generating synthetic polynucleotides which encode lysyl oxidase or lysyl oxidase-like proteins. Synthetic polynucleotides are nucleotide sequences which have been specifically designed for high level expression of lysyl oxidase or lysyl oxidase-like proteins in particular host cells.

Synthetic polynucleotides of the invention are generated by introducing specific nucleotide mutations at specific positions with respect to a particular codon, so as to replace codons which are rarely used, with codons which are frequently used by the particular host. Typically the mutation will not encode an amino acid To this extent, the nucleotide mutations within the synthetic polynucleotide are silent mutations. The number of codons corresponding to each amino acid residue is shown in Table 2. As a result of the degeneracy of the genetic code it is recognised that for some amino acid residues, silent mutations may be introduced at any one of nucleotide positions 1, 2 or 3 of a particular codon. It is further recognised that for some amino acid residues, for example cysteine, arginine, glutamine, few silent mutations may be introduced. particular methionine and tryptophan residues are both generally encoded by a single codon. Nonetheless, the present inventor anticipates that, as a result of the known low stringency codon-anti-codon interaction at nucleotide position 3 which gives rise to the "wobble effect", (Crick, F. et al. 1966) further silent mutations may be introduced at position 3, so as to increase the

number of codons for a particular amino acid residue which may be selected by a particular host cell.

The Genetic Code						
First Position (5' end)	tion Second Position					
	U .	С	A	G		
ט	PHE LEU LEU	SER SER SER SER	TYR TYR Stop Stop	CYS CYS Stop TRP	U C A G	
С	LEU LEU LEU	PRO PRO PRO PRO	HIS HIS GLN GLN	ARG ARG ARG ARG	U C A G	
· A	ILE ILE ILE MET	THR THR THR THR	ASN ASN LYS LYS	SER SER ARG ARG	U C A G	
G	VAL VAL VAL	ALA ALA ALA ALA	ASP ASP GLU GLU	GLY GLY GLY GLY	U C A G	
Note:	Given the position of the bases in a codon, it is possible to find the corresponding amino acid. For example, the codon (5')AUG(3') on mRNA specifies methionine, whereas CAU specifies histidine. UAA, UAG and UGA are termination signals. AUG is part of the initiation signal, and it codes for internal methionines as well.					

The present invention also relates to synthetic polynucleotides encoding lysyl oxidase or lysyl oxidase-like proteins wherein the nucleotide sequence of the synthetic polynucleotide comprises mutations which encode amino acid mutations. The lysyl oxidase and lysyl oxidase-like proteins which contain amino acid sequence mutations are variants of lysyl oxidase or lysyl oxidase-like proteins and are also described herein as variants of the invention. As described below, variants of the invention retain the functional properties of lysyl oxidase and lysyl oxidase-like proteins, namely that they catalyse oxidation of amine groups including lysine. The

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amino acid sequence of lysyl oxidase or lysyl oxidaselike proteins may be mutated by addition, deletion,
substitution or combinations thereof of the encoding
nucleotide sequence and such altered molecules are
variants of the invention provided they retain the
required activity. In particular, the inventor has
recognised that the amino acid sequence of lysyl oxidase
or lysyl oxidase-like proteins may be altered without
significantly changing the specificity or efficacy of the
enzyme, so that the enzyme containing the amino acid
mutation is substantially the same as the enzyme
comprising the native amino acid sequence. The inventor
also recognises that the specific activity and substrate
specificity may be altered by mutation of lysyl oxidase
or lysyl oxidase-like protein amino acid sequence.

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Thus the present invention further provides amino acid sequence variants of lysyl oxidase and lysyl oxidase-like proteins. Variants of the invention retain the functional properties of lysyl oxidase and lysyl oxidase-like proteins, namely that they catalyse 20 oxidation of amine groups including lysine. Variants of the invention have amino acid sequences which are homologous with the amino acid sequence of lysyl oxidase or lysyl oxidase-like proteins. For the purposes of this description, "homology" between the amino acid sequence 25 of a lysyl oxidase or lysyl oxidase-like protein and that of a variant connotes a likeness short of identity, indicative of a derivation of one sequence from the In particular, an amino acid sequence is homologous to that of a lysyl oxidase or lysyl oxidase-30 like protein if the alignment of that amino acid sequence with the sequence of a lysyl oxidase or a lysyl oxidaselike protein reveals a similarity of about 65% over any 20 amino acid stretch or over any repetitive element of the molecules shorter than 20 amino acids in length. 35 Such a sequence comparison can be performed via known algorithims such as the one described by Lipman and Pearson (1985), which are readily implemented by

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computer. Similarity is observed between amino acids where those amino acids have a side chain which confers a similar chemical property in the same chemical environment. For example, threonine and serine are similar amino acids; aspartic acid and glutamic acid are similar amino acids; valine, leucine and isoleucine are similar amino acids etc. Thus an amino acid sequence may be considered homologous with the amino acid sequence of a lysyl oxidase or a lysyl oxidase-like protein because the alignment of those sequences reveals a similarity of 65%, although at each amino acid position in the aligned sequences, none of the residues are identical.

According to a first aspect of the present invention, there is provided a synthetic polynucleotide encoding an amino acid sequence selected from the group consisting of lysyl oxidases, lysyl oxidase-like proteins and variants thereof.

The lysyl oxidase may be a vertebrate including human, bovine, porcine, rat or mouse lysyl oxidase, or avian lysyl oxidase including chick lysyl oxidase. Preferably, the lysyl oxidase is a human lysyl oxidase. The nucleotide sequences of rat and mouse lysyl oxidase are reported in Trackman et al. (1991), porcine in Cronshaw et al. (1993), mouse in Contente et al. (1993), and chick in Wu et al. (1992).

The lysyl oxidase-like proteins of the present invention are proteins which exhibit an amine oxidation catalysing function.

The variants of lysyl oxidases and lysyl oxidase-like proteins in accordance with the present invention may be truncated forms of the lysyl oxidase or lysyl oxidase-like proteins wherein the truncated forms retain the functional properties of lysyl oxidase and lysyl oxidase-like proteins, in particular the ability to catalyse the oxidation of amine groups, including lysine. In toto, the truncated forms are typically greater than 20 amino acids in length.

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The variants of the invention also include molecules that comprise a particular sequence of amino acids or a particular domain of a lysyl oxidase or lysyl oxidase-like protein wherein the sequence of amino acids or domain retain the functional properties of lysyl oxidase and lysyl oxidase-like proteins, in particular the ability to catalyse the oxidation of amine groups, including lysine.

Synthetic polynucleotides according to the invention are generated by mutating the native nucleotide sequence encoding a lysyl oxidase or lysyl oxidase-like protein or variant so that:

- a) all or some of the codons which hamper expression in the expression system in which the polynucleotide is to be expressed, are replaced with codons more favourable for expression in the expression system; and
 - b) the altered sequence encodes a protein or polypeptide selected from the group consisting of lysyl oxidases, lysyl oxidase-like proteins and variants thereof.

In one preferred embodiment the synthetic polynucleotide comprises a nucleotide sequence which encodes lysyl oxidase wherein the nucleotide sequence has a CAI score of at least 0.3 in a particular host cell. More preferably the synthetic polynucleotide comprises a nucleotide sequence which encodes human lysyl oxidase, wherein the synthetic polynucleotide has a CAI of at least 0.3 in E. coli. Most preferably the synthetic polynucleotide comprises all or part of the nucleotide sequence shown in Figure 2 (SEQ ID NO:3), starting at residue 141 (Met) or 144(Asp). In the sequence shown in Figure 2 (SEQ ID NO :3) the N-terminal encoded amino acid is serine. The presence of the serine residue is the consequence of the insertion of a BamHI restriction site at this position. It will be readily recognised that this serine residue could be deleted or replaced by an alternative residue related to the restriction site

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present at the N-terminal end of the construct. Similarly, at the 3' end of the construct, additional stop codons are inserted compared to the parent sequence shown in Figure 1 (SEQ ID NO: 1). These stop codons are a feature of this particular construct.

With respect to the sequence of the lysyl oxidase molecule, the first 21 amino acids of the sequence shown in Figure 1 (SEQ ID NO:2) are amino acids of the presequence. The residues from this position and up to and including residue 143 comprise the pro-sequence. Thus the mature form of the lysyl oxidase commences at residue 144. These different constructs are encompassed within the synthetic polynucleotides and encoded products of the invention.

In another preferred embodiment, at least 50% of codons in a particular nucleotide sequence which may be altered so as to reflect frequent codon usage in a particular host, are selected and altered. More preferably at least 60% are altered. Most preferably at least 70% are altered.

In another preferred embodiment at least 50% of codons which encode a particular amino acid are altered, so as to reflect frequent codon usage in a particular host. More preferably at least 60% are altered. Most preferably at least 70% are altered.

Modifications to codon usage to provide enhanced expression for a variety of hosts are discussed in:

Zhang et al. (1991), for E. coli, yeast, fruit fly and primates where codon usage tables are provided;

Newgard et al. (1986), for mammals; and

Newgard et al. (1986), for mammals; and
Murray et al. (1989), for plants. Preferred codon usages
are indicated in these publications.

It is important to recognise that even where human originating nucleotide sequences are to be expressed in human cell lines, or other mammalian cell lines, that modifications to codon usage can still be beneficial to producing enhanced levels of the protein of interest. The same applies to expression of other lysyl oxidase and

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lysyl oxidase like proteins in cell lines derived from their originating species or other mammalian or avian species.

In another preferred embodiment, all or part of the 5' or 3' untranslated region, or intronic regions of the synthetic nucleotide sequence encoding the lysyl oxidase or lysyl oxidase-like protein or variant thereof, are deleted.

In another preferred embodiment, all or part of the regions encoding the signal peptide, or regions encoding the pro-peptide, of the synthetic nucleotide sequence encoding the lysyl oxidase or lysyl oxidase-like protein or variant thereof, are deleted.

More preferably, all or part, of the 5' or 3'
untranslated region, and/or intronic regions, and regions
encoding the signal peptide, and/or regions encoding the
pro-peptide, of the synthetic polynucleotide encoding the
lysyl oxidase or lysyl oxidase-like protein or variant
thereof are deleted.

It will be recognised that the strategies for specific synthetic gene construction to yield enhanced expression of lysyl oxidases described in detail herein, can readily be adapted to the synthesis of other lysyl oxidases, lysyl oxidase-like proteins and variants thereof not specifically exemplified here, in light of the significant degree of homology between these molecules and in light of the fact that the present inventor has established that these proteins can be expressed in functional form in significant amounts from particular host cells using synthetic gene constructs with enhanced codon usage.

In another preferred embodiment the synthetic polynucleotide comprises a nucleotide sequence encoding lysyl oxidase, lysyl oxidase-like protein or a variant thereof and a further nucleotide sequence, wherein the nucleotide sequence encoding lysyl oxidase, lysyl oxidase-like protein or a variant thereof and the further nucleotide sequence are chemically linked so as to encode

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a single open reading frame. The further nucleotide sequence is preferably chemically linked to the 5' end of

the nucleotide sequence encoding lysyl oxidase, lysyl

oxidase-like protein or a variant thereof.

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In another preferred embodiment, the further nucleotide sequence encodes all of part of a polypeptide which causes the lysyl oxidase, lysyl oxidase-like protein or variant thereof to be either secreted to the extracellular environment, expressed as a cell surface periplasmic protein, or expressed in the intracellular environment. Preferably the further nucleotide sequence encodes all or part of glutathione S-transferase.

In another preferred embodiment the further nucleotide sequence encodes additional residues such as an N-terminal methionine or formyl-methionine.

In a preferred form of the invention the synthetic polynucleotide is unfused.

In another preferred embodiment, the synthetic polynucleotide comprises a fragment of the nucleotide sequence shown in Figure 2, wherein the fragment of the nucleotide sequence encodes a protein which retains the enzymatic activity of lysyl oxidase, lysyl oxidase-like protein or a variant thereof. Preferably the synthetic polynucleotide comprises the sequence shown in Figure 2 and commencing at the codon encoding residue 141. another preferred embodiment the synthetic polynucleotide comprises the sequence shown in Figure 2 and commences at or near the codon encoding residue 144.

In a further preferred embodiment, the synthetic 30 polynucleotide comprises the nucleotide sequence shown in Figure 2 (SEO ID NO:3), or a fragment thereof, in which only some of the nucleotide mutations have been introduced. Preferably, the synthetic polynucleotide comprises the nucleotide sequence shown in Figure 2 (SEQ ID NO:3), or a fragment thereof, and at least 50% of the 35 nucleotide mutations have been introduced. preferably, the synthetic polynucleotide comprises the nucleotide sequence shown in Figure 2 (SEQ ID NO:3) in

which the number of nucleotide mutations which have been introduced produce a CAI score for the synthetic polynucleotide in *E. coli* of at least 0.30.

The production of the synthetic polynucleotides of the invention, which is described in detail in the accompanying Examples, can be via the techniques of organic synthesis or recombinant DNA technology, or a combination of both techniques.

The synthetic polynucleotides described in the accompanying Examples were designed to make them amenable to cloning and expression in fused and unfused forms, and for intracellular and secreted production.

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Where the synthetic polynucleotide is prepared from assembled oligonucleotides it is preferred to incorporate restriction sites in the sequence to facilitate assembly of the polynucleotide.

Restriction sites incorporated in the polynucleotide sequence are also useful for:

- facilitating subcloning of manageable blocks for
 sequence confirmation;
 - 2. providing sites for later introduction of modifications to the polynucleotide as insertions, deletions or base changes;
- 3. facilitating confirmation of correct polynucleotide assembly by restriction endonuclease digestion.

The synthetic polynucleotides encoding the variants of the invention can be produced by the techniques of site-directed mutagenesis or random mutagenesis. These techniques allow one to determine the effect of mutation at particular positions in the synthetic polynucleotide sequence on the amine oxidation catalysing properties of the encoded variant of the invention.

In one preferred embodiment, the variants of the invention are produced by the technique of site-directed mutagenesis using oligonucleotides, which comprises the following steps:

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synthesis of an oligonucleotide with a sequence that contains the desired nucleotide mutation;

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hybridizing the oligonucleotide to a template nucleotide sequence encoding lysyl oxidase or a lysyl oxidase-like protein; and

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using a DNA polymerase to extend from the 3' end of the oligonucleotide in the 5' to 3' direction.

Another approach which is particularly suited to situations where the synthetic polynucleotide has been prepared from oligonucleotide blocks bounded by restriction sites is cassette mutagenesis where entire restriction fragments are inserted, deleted or replaced.

According to a second aspect of the present invention there is provided a recombinant nucleic acid molecule comprising a synthetic polynucleotide of the first aspect. Preferably the recombinant nucleic acid molecule comprises vector nucleic acid.

The vectors of the invention include plasmids, phages, and phagemids. The vectors of the invention can be used in integrative expression systems or lytic or comparable expression systems.

Suitable vectors will generally contain origins of replication and control sequences which are derived from species compatible with the intended expression host.

- Typically these vectors include a promoter located upstream from the synthetic polynucleotide, together with a ribosome binding site if intended for prokaryotic expression, and a phenotypic selection gene such as one conferring antibiotic resistance or supplying an auxotrophic requirement. For production vectors, vectors which provide for enhanced stability through partitioning may be chosen. Where integrative vectors are used it is not necessary for the vector to have an origin of
- replication. Lytic and other comparable expression systems do not need to have those functions required for 35 maintenance of vectors in hosts.
 - For E. coli typical vectors include pBR322, pBluescript II SK+, pGEX-2T, pTrc99A, pET series vectors,

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particularly pET3d, (Studier et al. 1990) and derivatives of these vectors.

In one preferred embodiment, the vector comprises a synthetic polynucleotide of the first aspect, wherein the synthetic polynucleotide has a CAI score of at least 0.30 in a particular host cell. Preferably the vector comprises all or part of a synthetic polynucleotide sequence shown in Figure 2 (SEQ ID NO:3).

In another preferred embodiment, the vector comprises a synthetic polynucleotide of the first aspect wherein at least 50% of codons in a particular nucleotide sequence which may be altered so as to reflect frequent codon usage in a particular host, are selected and altered. More preferably at least 60% are altered. Most preferably at least 70% are altered.

In another preferred embodiment the vector comprises a synthetic polynucleotide of the first aspect wherein at least 50% of codons which encode a particular amino acid are altered, so as to reflect frequent codon usage in a particular host. More preferably at least 60% are altered. Most preferably at least 70% are altered.

In another preferred embodiment the vector comprises a synthetic polynucleotide of the first aspect wherein all or part of the 5' or 3' untranslated region, and/or intronic regions is deleted.

In another preferred embodiment the vector comprises a synthetic polynucleotide of the first aspect wherein all or part of the regions encoding the signal peptide, and/or regions encoding the pro-peptide is deleted.

In another preferred embodiment the vector comprises a synthetic polynucleotide of the first aspect wherein all or part, of the 5' or 3' untranslated region, and/or intronic regions, and/or all or part of the regions encoding the signal peptide, and or regions encoding the pro-peptide is deleted.

In another preferred embodiment, the recombinant DNA molecule comprises a synthetic polynucleotide of the first aspect, wherein the synthetic polynucleotide

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sequence is chemically linked to a further nucleotide sequence so as to encode a single open reading frame for the synthetic polynucleotide and the further nucleotide sequence. Preferably, the recombinant DNA molecule comprises a synthetic polynucleotide and the further nucleotide sequence wherein the further nucleotide sequence is chemically linked to the 5' end of the synthetic polynucleotide.

In another preferred embodiment, the recombinant DNA molecule comprises a synthetic polynucleotide, wherein the synthetic polynucleotide comprises all or part of the nucleotide sequence shown in Figure 2 (SEQ ID NO:2). Preferably, the recombinant DNA molecule comprises a synthetic polynucleotide sequence having all or part of the sequence shown in Figure 2 (SEQ ID NO:3) wherein only some of the indicated nucleotide mutations have been introduced.

According to a third aspect of the present invention there is provided a host cell transformed with a recombinant DNA molecule of the second aspect of the invention.

The host cells may be prokaryotic cells including bacteria, and may also include yeasts, insect cells and other eukaryotic cells or whole organisms.

In a preferred embodiment, the host is an *E. coli* strain including *E. coli* B strain derivatives (Studier *et al.* 1990), and *E. coli* K12 derivatives such as NM522 (Gough and Murray, 1983) and XL1-Blue (Bullock *et al.* 1987).

In another preferred embodiment, the host is S. cerivisiae or Pichia.

According to a fourth aspect of the present invention there is provided an expression product, wherein the expression product is derived from a host cell of the third aspect, the expression product comprising an amino acid sequence selected from the group consisting of lysyl oxidase, lysyl oxidase-like proteins and variants thereof.

In one preferred embodiment, the expression product comprises the amino acid sequence shown in Figure 2 (SEQ ID NO:4) and is encoded by a synthetic polynucleotide which comprises the nucleotide sequence shown in (SEQ ID NO:3).

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In an alternative embodiment, the expression product comprises a fragment of the amino acid sequence shown in Figure 2 (SEQ ID NO:4) which retains amine oxidation catalysing properties, and is encoded by a synthetic polynucleotide comprising the corresponding nucleotide sequence shown in Figure 2 (SEQ ID NO:3). Preferably the expression product comprises an amino acid sequence having an N-terminus at amino acid residue position 141 (Met) of Figure 2 (SEQ ID NO:4) or results in an N-terminal amino acid residue position 144 (Asp) of Figure 2, (SEQ ID NO:4) and is encoded by a synthetic polynucleotide comprising the corresponding nucleotide sequence shown in Figure 2 (SEQ ID NO:3).

In another preferred embodiment, the expression product is encoded by a synthetic polynucleotide 20 comprising the nucleotide sequence shown in Figure 2 (SEQ ID NO : 3) in which only some of the nucleotide mutations have been introduced. Preferably the expression product is encoded by a synthetic polynucleotide comprising all 25 or part of the nucleotide sequence shown in Figure 2 (SEQ ID NO:3) and at least 50% of the nucleotide mutations relative to Figure 1 have been introduced. preferably the expression product is encoded by a synthetic polynucleotide comprising all or part of the nucleotide sequence shown in Figure 2 (SEQ ID NO:3) in 30 which the number of nucleotide mutations which have been introduced produce a CAI score for the synthetic polynucleotide in E. coli of at least 0.30.

The expression products of the fourth aspect may be fused expression products which include all or part of a protein encoded by the vector in peptide linkage with the expression product. They may also include, for example, an N-terminal methionine or other additional residues

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which do not permanently impair the catalytic properties of the product.

In a further preferred embodiment, the expression products of the invention are fusion products which comprise an amino acid sequence selected from the group consisting of lysyl oxidase, lysyl oxidase-like proteins and variants thereof which are encoded by a synthetic polynucleotide, wherein the amino acid sequence is chemically linked to a further amino acid sequence.

Typically, the further amino acid sequence is chemically linked to the N-terminus of the expression product. Preferably the further amino acid sequence allows purification of the fusion product. Most preferably the further amino acid sequence comprises an amino acid sequence encoding glutathione S- transferase. The further amino acid sequence may be chosen in order to cause the fusion product to be secreted to the extracellular evironment, expressed as a cell membrane protein or expressed in the intracellular environment.

The expressed fusion products may subsequently be treated to remove the further amino acid sequence from the amino acid sequence selected from the group consisting of lysyl oxidase, lysyl oxidase-like proteins and variants thereof. Preferably, the treatment is via chemical cleavage. In another more preferred embodiment, the treatment is via protease digestion. Still more preferably, the treatment is effected by endogenous host cell secretion machinery, for example yeast cell secretion machinery.

Non-fused systems include the introduction of or use of a pre-existing methionine codon. An example of this is the use of pET3a and pET3d in *E. coli*.

In a further embodiment, the invention provides a pharmaceutical composition comprising an expression product of the invention and a pharmaceutically acceptable carrier, excipient or diluent. The pharmaceutical compositions are prepared and tested according to standard pharmaceutical practise. The

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support for their efficacy is provided by the assays in the Examples.

According to a fifth aspect of the invention there is provided a method for producing an expression product of the invention, the method comprising the following steps:

- (i) providing a host cell of the third aspect of the invention.
- (ii) maintaining the host cell in conditions
 suitable for expression of the synthetic polynucleotide;
 and
 - (iii) isolating the expression product.

In one preferred embodiment the expression product is produced by expression of the synthetic polynucleotide in *E. coli*. The invention extends to the use of other host cells which are capable of producing the expression product by expression of the synthetic polynucleotides designed for use in *E. coli*.

In another preferred embodiment, the expression product is produced by expression of a synthetic polynucleotide of the invention in other microbial expression systems including prokaryotic expression systems such as bacterial expression systems, and also including insect cell expression systems, and expression systems involving other eukaryotic cells or whole organisms. Again synthetic polynucleotides which are designed for use in a particular host can also be used in other cells which provide high expression of that sequence.

As disclosed in the Examples, the inventor has

determined that it is useful to add Cu to the expression
product which has been purified from the inclusion bodies
of E. coli or similar organisms, so that the expression
product can fold into a functional conformation which has
the enzymatic activity of lysyl oxidase or lysyl oxidaselike proteins. The role of Cu in solubilising lysyl
oxidase protein was not apparent until the present
invention was carried out. Indeed based on earlier
reported results which show that inactive apoenzyme can

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be reactivated by the addition of Cu it would have been expected that re-folding could occur without Cu. The present inventor initially attempted to refold protein in the absence of Cu and failed to generate soluble protein. It was only in subsequent experiments where Cu was added that refolding in this environment succeeded.

Further, should over expressed lysyl oxidase lack copper after its production in bacteria, the incorporation of copper into the recombinant protein would be possible by adapting the previously reported protocol (Gacheru et al. 1990) used for conversion of the apoenzyme of natural lysyl oxidase into the metal containing form.

In addition, it would be desirable to add oxidising substances that assist in the production of the organic cofactor if this is not formed by the enzyme under the prevailing conditions without assistance. The most important additional substance is oxygen in solution.

It is proposed here that generation of the active enzyme in vitro is a time-dependent process involving the introduction of copper to the overexpressed polypeptide, followed by oxidation of the protein to generate the covalently bound quinone. This is despite the absence of a Asn-Tyr-Asp/Glu consensus sequence from lysyl oxidase presumed to be important to this type of process (Tanizawa 1995).

The expression product may be exposed to chemical agents to enhance the stability and activity of the molecule. The chemical agents include various concentrations of urea or glycerol combined with storage at -20°C.

In one preferred form the expression product is produced in the form of inclusion bodies which are harvested from the transformed host.

As described above, the inventor recognised a variety of uses for lysyl oxidase, lysyl oxidase-like proteins and variants thereof although these uses require significant amounts of protein in functional

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conformation. As the inventor has for the first time produce significant amounts of enzyme in functional conformation, the following uses, as described below can now be realised.

The invention provides a method of cross-linking molecules that contain primary amines, wherein the molecules are associated with a cell membrane or extracellular environment, comprising contacting the primary amine with an expression product of the invention.

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As described in the Examples, the inventor now recognises that the expression products of the invention have specific activity for primary amine groups in peptide and non-peptide substrates. Furthermore, as described in the Examples, as the present inventor has, for the first time, observed cross-linking of both peptide and non-peptide substrates in vitro, using the expression product of the invention, the inventor envisages that the expression products of the invention will be useful for cross-linking a wide range of molecules such as peptides and polypeptides, non-peptide chemical polymers which contain primary amine groups and combinations thereof. In particular, the present inventor envisages that the expression product of the invention will be widely applicable to methods of treatment in vivo, in particular where those methods of treatment require cross-linking of elastin or collagen. For this purpose the expression product can be formulated in suitable carriers, buffers and other conventional delivery systems.

In one preferred embodiment, the invention provides a method of cross-linking molecules that contain primary amines, wherein the molecule is a peptide or polypeptide molecule and the primary amine is associated with an amino acid side chain. The method comprises contacting the peptide or polypeptide molecule with an expression product of the invention under conditions suitable to cross-link the molecules. Preferably, the peptide or

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polypeptide comprises the amino acid sequence of tropoelastin or a fragment thereof. In another preferred embodiment, the peptide or polypeptide comprises the amino acid sequence of collagen or a fragment thereof. It can also comprise both.

In another preferred embodiment the invention provides a method of cross-linking molecules that contain primary amines, wherein the the molecule is a non-peptide molecule. The method comprises contacting the non-peptide molecule with an expression product of the invention under conditions suitable to cross-link the molecules.

The present inventor also recognises that the $^{ au}$ expression products of the invention would find wide application in accelerating the rate of wound healing. In particular the inventor has recognised that lysyl oxidase, lysyl oxidase-like proteins and variants thereof may be used to accelerate wound healing. Wound healing may be accelerated by administering an expression product of the invention to the wound. Alternatively, an expression product of the invention may be included in a matrix including an elastin, tropoelastin and/or collagen based or amine containing matrix which is itself applied to the wound. In both methods, wound healing is accelerated by cross-linking of molecules which contain primary amines, and the cross-linking is mediated by the expression product of the invention. For this purpose the expression product can be formulated in suitable carriers, buffers and other conventional delivery systems.

Thus the invention provides a method of accelerating the rate of wound healing, the method comprising the step of administering an expression product of the invention to a wound under conditions which are suitable for crosslinking of molecules containing primary amines.

Preferably, an expression product of the invention is administered to the wound by direct application of the product to the wound. More preferably the product is

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administered to the wound by first including the product in a matrix which is itself administered to the wound. Most preferably, the product is administered to the wound by application of a pharmaceutical formulation comprising the expression product of the invention and a pharmaceutically acceptable carrier, diluent or excipient.

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In another preferred embodiment, the invention relates to a method of deccelerating the rate of wound healing. The inventor recognises that there are 10 particular surgical procedures which require a delay in wound healing so as to achieve a desirable outcome. A lysyl oxidase-mediated delay in wound healing is based on the ability to inhibit the oxidation of primary amines 15 and therefore the ability to inhibit the cross-linking of particular molecules, where that cross-linking is essential to wound healing. Lysyl oxidase inhibitors are known, for example BAPN, although these reagents are typically noxious and therefor unsuitable for use in methods of treatment. The inventor envisages that a 20 delay in wound healing which is mediated by the inhibition of oxidation of primary amines can be achieved by developing novel reagents which have specific antilysyl oxidase acitvities and which have no side-effects. Particularly suitable inhibitors of lysyl oxidase 25 activities are anti-lysyl oxidase antibodies. immunological reagents may be generated by raising antibodies to the expression products of the invention. The methods for the production of monoclonal antibodies or polyclonal antibodies are within the knowledge of the 30 skilled addressee and are described in detail in standard immunological texts. Furthermore, the skilled addressee can readily determine those antibodies with anti-lysyl oxidase activities, in particular the ability to inhibit the oxidation of primary amines by performing screening 35 of those antibodies according to the benzylamine substrate assay, tropoelastin substrate assay or

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tritiated tropoelastin substrate assay, as described in the accompanying Examples.

Thus, the invention provides a method of deccelerating the rate of wound healing, the method comprising the step of administering antibodies which are raised against the expression product of the invention, to a wound under conditions which allow the antibodies to substantially inhibit the oxidation of primary amines.

In another aspect embodiment, the invention relates to novel reagents for modifying the rate of wound healing. In particular, the inventor recognises that the expression products of the invention can be used to link chemical species to heterologous surfaces to substantially promote or substantially inhibit the rate of wound healing. The chemical species of interest generally include those species which contain a primary amine and which can be linked by oxidation of the primary amine to a heterologous surface, but more specifically comprise those chemical species selected from the group consisting of immunomodulatory substances including growth factors and cytokines, and substances which modify inflammatory responses or the coagulation cascade and also antibiotic substances. The heterologous surface may be a matrix comprising a complex of biomolecules, in particular elastin or tropoelastin, or a complex of nonpeptide molecules. The expression product of the invention may be used to form a linkage between two chemical species which contain primary amine groups. Thus the invention has particular application to the attachment of an active ingredient to a carrier molecule.

Thus the invention provides novel reagents and formulations thereof for use in modifying the rate of wound healing, wherein the reagents are generated by oxidising a primary amine contained in a chemical species so as to link the chemical species to a heterologous surface.

In another preferred embodiment, the invention relates to the use of an expression product of the

invention in the treatment or prevention of abnormal fibrosis. Lysyl oxidase has an important functional role in fibrosis and is also believed to be involved in the abnormal process of fibrosis, in particular in the fibrotic liver (Kagan H. M. 1994). The present inventor envisages the use of the expression products of the invention in promoting fibrosis so as to achieve a normal physiological outcome and inhibiting fibrosis where an abnormal outcome is to be avoided. Fibrosis may be promoted by increasing the levels of available lysyl 10 oxidase or by increasing the specific activity of the enzyme. For example fibrosis may be promoted by administering an expression product of the invention to a particular site so as to promote fibrosis at that site. Fibrosis may also be promoted by administering anti-lysyl 15 oxidase antibodies which augment or enhance the specific activity of the enzyme, for example by interfering with aggregation of the protein. Fibrosis may be inhibited by inhibiting or reducing the ability of the lysyl oxidase to oxidise primary amine residues. The inventor envisages that fibrosis, as mediated through the role of lysyl oxidase in the oxidation of primary amines, could be inhibited by the use of antibodies which inhibit the oxidation reaction. As described above, the methods for the production of monoclonal antibodies or polyclonal antibodies are within the knowledge of the skilled Furthermore, the skilled addressee can addressee.

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readily determine those antibodies with anti-lysyl oxidase activities which either inhibit or enhance lysyl oxidase activity, in particular the ability to inhibit or 30 enhance the oxidation of primary amines by performing screening assays according to the benzylamine substrate assay, tropoelastin substrate assay or tritiated tropoelastin substrate assay, as described in the 35 accompanying Examples.

As described above, three separate assays are disclosed in the accompanying Examples for measuring levels of lysyl oxidase activity. The inventor

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recognises that the expression products of the invention could be used in a particular method so as to screen agents for the ability to inhibit or enhance lysyl oxidase acitivity. Agents which are identified as having inhibitory or enhancing effects on the specific activity of lysyl oxidase would be desirable for use in specific treatments, including wound healing or fibrosis, as described above.

Thus in a preferred embodiment, the invention provides a method for screening agents which substantially enhance or substantially inhibit the specific activity of lysyl oxidase or lysyl oxidase-like proteins, the method comprising the step of contacting the agent with an expression product of the invention and measuring the specific activity of the expression product.

To the extent that lysyl oxidase and lysyl oxidaselike proteins are involved in the oxidation of primary amines, the inventor recognises that the expression products of the invention are useful in assays for 20 detecting substrates which contain primary amines. expression products of the invention are also useful to quantify amounts of protein or non-protein substrates wherein those substrates contain a known number of primary amine moieties. The expression products of the 25 invention are particularly suited to such an assay as the expression product is derived from a pure source and will therefore specifically detect primary amine groups. Furthermore, as the expression products of the invention may be variants of lysyl oxidase or a lysyl oxidase-like 30 protein, the inventor recognises that molecules associated with a particular primary amine (for example a primary amine group displayed in a specific manner) may be specifically detected over other molecules in the 35 sample which contain a primary amine group. The assay could be performed according to any of the three substrate assays detailed in the accompanying Examples.

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Thus in a preferred embodiment, the invention relates to a method for detecting a protein or nonprotein substrate in a sample, the method comprising contacting the sample with an expression product of the invention and detecting oxidation of any primary amine moieties in the sample.

In another preferred embodiment, the invention relates to a method for quantifying the amount of a protein or non-protein substrate in a sample, wherein the number of primary amine moieties per substrate molecule is known, the method comprising contacting the sample with an expression product of the invention and detecting the amount of oxidation of any primary amine moieties in the sample.

In one aspect, the lysyl oxidase, lysyl oxidase-like proteins and variants thereof may be used to cross-link a primary amine containing molecule such as tropoelastin or collagen to the surface, wherein the solid surface has primary amine groups which are available for crosslinking. In this aspect, the invention provides the use 20 of lysyl oxidase, lysyl oxidase-like proteins and variants thereof as an agents for forming coatings on solid surfaces. The present inventor envisages that those surfaces which have primary amine groups available for cross-linking will be useful for the manufacture of synthetic tissues for example, blood vessels.

In one embodiment, it is envisaged that surface or devices could be dipped into solutions of tropoelastin and/or collagen in the presence of a lysyl oxidase, lysyl oxidase-like protein or variant of the invention.

In another embodiment, a cross-linked material could be preformed and then cross-linked to available primary amine groups on a surface, using a lysyl oxidase, lysyl oxidase-like protein or variant of the invention.

The cross-linked molecule may comprise collagen and/or tropoelastin and/or other cross-linkable primary amine containing molecules.

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In another aspect, the invention provides a tissue glue, which is of use in assisting wound healing. glue can be used as a replacement for sutures. The glue comprises an expression product of the invention and may 5 additionally comprise at least one cross-linkable primary amine material, for example, tropoelastin and/or The glue may also be used to repair incisions collagen. in for instance, surgical applications and other body tissue defects. The glue may be applied by mixing the primary amine containing material and expression product and applying to the site.

Other uses for the expression products of the invention include the preparation of sheets or films where the product can be used as a reinforcement as well as for its biological properties. Applications could include repair of anastomoses, repair of defects in the The expression products can also be used to prepare solid structures such as beads which may find application in tissue bulking applications such as in the urethra, oesaphagus or in dermatological applications where primary amine containing substances are desirably deposited in cross-linked form.

The expression products of the invention are also useful as agents for:

The routes envisaged for the administration of 25 this form of cancer therapy include gene and protein therapies in which the gene or protein is administered in accordance with protocols already developed with respect to other molecules proposed for use in gene or protein therapy. 30 application category it is proposed that extracellular application of lysyl oxidase either alone or as part of a delivered complex, could reduce the growth and behaviour of cancer and other cells, including but not limited to ras-transformed 35 Lysyl oxidase administration has the potential to display the anti-cancer phenotypic

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potential of drugs such as azatyrosine (Krzyzosiak et al., 1992; Contente et al., 1990).

- Regulating cell growth. Increased lysyl oxidase may be considered to mimic some of the functions mediated by transforming-growth-factor-beta 1 (Shibanuma et al., 1993).
- Glue to join primary amine materials e.g. in surgical or veterinary applications. The expression product can be formulated in suitable carriers, buffers and other conventional delivery systems. This includes natural materials, synthetic materials and composites.
 - Oxidising accessible lysines (and other amines) to modify behaviour of those molecular components.
 - Modified lysyl oxidase to perform further reactions.

BRIEF DESCRIPTION OF ACCOMPANYING DRAWINGS

Figure 1: Entire coding sequence (SEQ ID NO:1) for an amino acid sequence (SEQ ID NO:2) of a secreted form of human lysyl oxidase. A potential signal sequence (1 to 21 aa) is underlined (Mariani et al., 1992; Hamainen et al., 1991; Svinarich et al. 1992).

Figure 2: DNA sequence (SEQ ID NO: 3) of a synthetic polynucleotide SHLOX encoding the amino acid sequence (SEQ ID NO: 4) of a lysyl oxidase. Restriction sites at the boundaries of each block are identified;

Figure 3: Comparison of the synthetic sequence SHLOX with the natural sequence (GenBank HUMLOX coordinates).

HUMLOX is on the upper line and SHLOX on the lower line, in each case;

Figure 4: Sequences of synthetic oligonucleotides for SHLOX construction (5' is at the left of each sequence, and 'p' denotes phosphate). (SEQ ID NOS:5-30)

Figure 5: Measure of lysyl oxidase activity as indicated by the production of tritiated water (TCA precipitated counts).

Figure 6: Measure of lysyl oxidase activity as indicated by the production of tritiated water.

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BEST METHOD OF CARRYING OUT THE INVENTION

Design of a synthetic lysyl oxidase gene

Human lysyl oxidase is typically made in human cells as a longer form, which is proteolytically processed to generate a shorter protein secreted into the extracellular space. The amino acid sequence has been compiled from data from different sources: complete sequence data which was available for cDNA encoding human lysyl oxidase (Hamalainen et al., 1991; Mariani et al., 1992), as well as partial human genome sequence (Svinarich et al., 1992). Analytical comparison of the cDNA sequences showed a discrepancy in amino acid composition which required elucidation before the synthetic gene could be made, namely Ala136(GCT) was in conflict with Arg136(CGT) of Hamalainen et al., (1991). 15 The present inventor dealt with this by considering the genomic data, and analysing homologous regions of lysyl oxidase from different species, including rat, mouse, bovine and chicken. The choice made was later confirmed (Hamalainen et al., 1993) supporting the correction that had been made in generating the synthetic gene. synthetic gene was designed with strategically placed restriction sites to facilitate conversion to novel derivatives and counterparts of lysyl oxidase and lysyl oxidase-related forms (Kim et al., 1995). The synthetic 25 gene was designed with a codon usage pattern favourable to high level expression in E. coli (Figure 2).

As lysyl oxidase is a secreted protein its cDNA derived sequence encodes a region for a signal peptide, which would not be useful for generating a functional enzyme in E. coli but rather for directing eventual transfer of the newly translated protein from inside the cell into the extracellular matrix. Furthermore the secreted pro-lysyl oxidase was thought to be cleaved by extracellular protease(s) to process the mature protein, yet it was not known if the pre-cleaved protein was active. There were doubts regarding the precise

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boundaries of the presumably removed signal peptide and propeptide ("leading peptide"). The likely leading peptide sequence in the N-terminal part of the lysyl oxidase sequence was determined by computer analysis and from the definitions provided by others (Trackman et al., 1990; Hamalainen et al., 1991) and was excluded from gene synthesis. Subsequent to this, further modifications were made to adjust for varied N-terminal sequences, various candidate signal sequences and eventually a better estimate of the leading peptide sequence, to generate derivatives of SHLOX, as detailed below.

Initial gene construction and cloning was conducted as described by Martin et al. (1995). The lysyl oxidase sequence of Figure 1 was backtranslated according to a table of average codon frequencies for a group of highly expressed E. coli genes (GCG package version 7-UNIX using CodonFrequency and GenRunData: ecohighcod). The resulting codon sequence lacked rare codons.

Modifications were made to the refined sequence to allow

for convenient construction, cloning and analysis by adding short artificial sequences to each end of the codon-optimized gene to provide BamHI-compatible cohesive termini, an NcoI site near the 5' end of the gene, and three stop codons at the end of the open reading frame.

Restriction sites were introduced into the synthetic coding sequence at 270 - 300 base pair intervals, dividing the gene into 4 sections of a manageable size for individual assembly, cloning and sequence analysis.

Each section was subdivided into pairs of

complementary oligonucleotides. Internal junctions
between oligonucleotide pairs were delineated by
nucleotide overlaps, whilst the external junctions fall
at the restriction enzyme cleavage points. The codonoptimized sequence and restriction sites are shown in

Figure 2, together with the encoded amino acid sequence.
Oligonucleotide sequences and designations are shown in
Figure 4.

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Oligonucleotide pairs 2 to 6 were made by the Sydney University Macromolecular Analysis Centre, Australia and subsequently purified by denaturing polyacrylamide gel electrophoresis (PAGE) as follows. Oligonucleotides (>85-mers) were run on PAGE containing 7M urea and 8% to 5 10% acrylamide (acrylamide:bisacrylamide 19:1). Samples were heated to 75°C for 3 minutes before loading. Tracking dye (0.05% bromphenol blue, 0.05% xylene cyanole FF in deionized formamide) was loaded into an adjacent lane and electrophoresis conducted at constant power of 10 25 W (ca. 300 V) until the bromphenol marker was within lcm of the base of the gel. Product bands were visualized by UV-shadowing over a fluorescent TLC plate. Gel fragments containing purified oligonucleotides were excised and transferred to microcentrifuge tubes, crushed 15 and soaked overnight at 60°C in $500\mu l$ elution buffer (0.3M sodium acetate pH7.0). A second extraction was performed with $400\mu l$ elution buffer, for 3 to 4 hours at 60°C and the supernatant combined with that of the first extraction. The total volume of the DNA-containing 20 solution was reduced to approximately $400\mu l$ by extraction with n-butanol and DNA was precipitated with 1ml 96% Purified oligonucleotides were collected by ethanol. centrifugation, redissolved in $20\,\mu l$ TE buffer and quantified by spectrophotometry. The final yield of 25 oligonucleotide was typically 5 to $30\mu g$. Oligonucleotides 1 and 7 to 13 were synthesised and purified by Macromolecular Resources, USA.

Complementary oligonucleotide pairs (1µg each, approximately 30pmol for 95-mer) were annealed in 10µl buffer containing 50mM Tris.HCl pH7.5, 10mM MgCl₂. The mixture was overlaid with 12µl paraffin oil, heated to 95°C and cooled to 16°C over 16h in a microprocessor-controlled heating block (Perkin Elmer Cetus Thermal Cycler). Annealed samples were transferred to clean microcentrifuge tubes and a small aliquot (1µl) withdrawn for analysis by agarose gel electrophoresis.

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For each block comprising three complementary oligonucleotide pairs (blocks 1-3), four separate reactions were set up (A+B, A+C, B+C and A+B+C). Each reaction contained 50mM Tris.HCl, 10mM MgCl₂, 1mM ATP,

5 3mM DTT, 3μl each of the appropriate annealed oligonucleotide pairs, 0.5μl (0.5U) T4 DNA ligase and Milli-Q water to a total volume of 10μl. All ingredients except ATP, DTT and T4 ligase (Boehringer Mannheim, Germany) were mixed and heated to 45°C for 5 minutes to

denature cohesive termini and then cooled to room temperature before addition of the remaining components. Ligation reactions were incubated overnight at 16°C and analysed on 2% LMT agarose gels, and ligated blocks were purified on preparative 2% LMT agarose gels with TAE

running_buffer. Product bands were identified under long-wave UV illumination with reference to known DNA size-standards (pBSIISK+ digested with either HaeIII or AluI) (Statagene, USA) and excised in the minimum possible volume of gel. DNA was recovered from LMT

agarose fragments using GELase or agarase. Block 4 was constructed similarly, except that four instead of three complementary oligonucleotide pairs participated in the ligation reaction. Purity and yield of recovered SHLOX blocks was assessed by analytical agarose electrophoresis alongside known DNA standards.

pBSIISK+ (Stratagene) DNA was digested with appropriate restriction enzymes (Figure 2). Inserts were purified at each stage by preparative gel electrophoresis. Plasmid DNA was isolated from agarose gels (Prep-A-Gene, Bio-Rad). Approximately 100ng (ca. 0.05pmol) purified plasmid fragment was added to 50ng (ca. 0.3pmol) synthetic block in 17µl buffer containing 50 mM Tris.HCl pH 7.5, 10mM MgCl₂ and the solution heated at 45°C for 5 minutes to denature cohesive termini. Upon cooling to room temperature, 2µl 10mM ATP, 30mM DTT and 1µl T4 DNA ligase (1U) were added and the reaction incubated overnight at 16°C. The mixture was heated for 10 minutes at 75°C to inactivate T4 ligase, and DNA was

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precipitated by adding $2.5\mu l$ 3M sodium acetate and $250\mu l$ ethanol to the $20\mu l$ volume of the ligation mixture. pellet was resuspended in a final volume of 10 μ l TE, pH 8.0. Of this solution, 1 to 2 μ l was used to electroporate 40µl XL1Blue competent cells (Bio-Rad electroporation Gene Pulser and Pulse Controller; 2500V/0.05sec). Electroporated cells were diluted with 1ml LB media immediately after the procedure, incubated 1 hour at 37°C and $100\mu l$ of this was spread on agar plates containing $100\mu g/ml$ ampicillin (LB plus ampicillin), 10 0.1mM IPTG and $80\mu g/ml$ X-gal. Recombinants were selected on the basis of a white bacterial colony phenotype, although some blue colonies were used as controls. Plasmid DNA was screened by restriction digestion and confirmed by sequencing. Plasmids containing the four 15 lysyl oxidase blocks were designated pSHLOX1-4.

All candidate clones were screened and the mutants discarded, unless they contained mutations of potential value in future SHLOX studies. Plasmids bearing correct sequences were grown in *E.coli* XL1Blue cells (Bullock et al 1987), purified by CsCl-ethidium bromide density gradient centrifugation and confirmed by sequencing in both directions.

In two separate reactions, adjacent pairs of gene blocks 1+2 and 3+4 were ligated in pBSIISK+. These were designed to give recombinants α and β , respectively. Each block was excised from the appropriate plasmid and purified through agarose gel electrophoresis. 50ng each synthetic block and 100ng appropriately restriction enzyme cleaved complementary-tailed pBSIISK+ digest were ligated in a total of 20μ l T4 ligase buffer in the presence of 1U T4 DNA ligase overnight at 16°C. Transformants were screened by the colony white phenotype, comparative electrophoretic mobility of the intact plasmids in 0.7% TBE agarose gel and by restriction digestion. After verification of the primary structure by sequencing, pSHLOX α and pSHLOX β were used for the final assembly of the pSHLOX gene.

pSHLOXα and pSHLOXß plasmids were cut and the appropriate fragments containing α and ß subassemblies of the intact gene of lysyl oxidase were purified from an agarose gel and ligated. Transformants were selected from LB plus ampicillin plates containing IPTG and X-gal. Correct constructs were confirmed by sequencing. The intact 1188bp synthetic gene encoding lysyl oxidase (SHLOX) was used for all further manipulations. The construct containing full length SHLOX (Figure 2) in pBSIISK+ was designated pSHLOXB.

The construction of SHLOX is exemplary of the methods used with other genes.

Construction of derivatives (including shorter forms) and placing in various E. coli vectors for production

15 <u>1. pGEXLOX-2T</u>

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The full length SHLOX gene was excised from pSHLOXB with BamHI and purified by gel electrophoresis and GELase (Epicentre Technologies, USA). Purified fragment (100ng) was ligated with 50ng pGEX-2T (Medos Co Pty Ltd) in 20µl total volume of ligase buffer using 1U of T4 DNA ligase. The vector DNA had been previously linearised with BamHI and treated with calf intestinal phosphatase (CIP; Boehringer Mannheim, Germany). Transformants were selected on LB+ampicillin plates, screened by restriction digestion and confirmed by DNA sequencing. The SHLOX gene cloned into pGEX-2T was designated pGEXLOX-2T.

2. pGEXLOX^{m1}

SalI and XhoI mutations were introduced (Clontech Transformer mutagenesis kit) into pGEXLOX-2T at positions

-13 to +23 and 421 to 442 of the SHLOX gene using two mutagenic primers (XhoI: CTG GTT CCG CGT GGA TCC CTC GAG CAG CCG CCG CGT; SalI: ATG GTT GGT GTC GAC CCG TAC A)

(Beckman Oligofax Service, Sydney). The 424 bp DNA fragment between the introduced SalI and XhoI sites was

excised from the 5'-terminal region of SHLOX of pGEXLOX-2T by cutting with SalI and XhoI and the remaining

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plasmid DNA was gel-purified. SalI and XhoI compatible ends were ligated. Apart from GST, the resulting construct pGEXLOX^{ml} was designed such that following thrombin cleavage of the expressed protein it would imitate a 'mature' SHLOX sequence, encoding a 30kDa protein with similar size and primary structure to the natural extracellular form of lysyl oxidase (Cronshaw et.al., 1995). Following sequence confirmation and further transformation, expression was in E.coli DH5α. The resulting expressed protein was designated GST-SHLOX.

3. p3aLml and p12aLml

pGEXLOX^{ml} was treated with BamHI to release the coding sequence SHLOX^{ml} (the truncated form of SHLOX). This was ligated, in each case, into the BamHI site of pET3a and pET12a plasmid vectors (Novagen Inc, USA) and the resulting constructs, p3aL^{ml} and p12aL^{ml} were transformed into E.coli HMS174 (Studier et al 1990). Transformants were plated and screened. Orientation of inserts was confirmed by restriction digestion and the regions around the cloning sites by sequencing. Following further transformation, expression was in E.coli BL21/DE3 (Studier et al 1990) cells for p3aL^{ml} and p12aL^{ml}. The resulting expressed proteins were designated 3aL^{ml} and 12aL^{ml}, respectively.

25 <u>4. p3DLox</u>

Full length SHLOX was gel-purified from pSHLOXB after BamHI digestion. pET3d vector was linearised with BamHI and dephosphorylated with CIP. Ligation of 100ng treated vector and 50ng purified SHLOX gene with T4 DNA ligase was conducted in 20µl volume at 16°C for 20h, and the ligation mixture was used to electroporate E.coli HMS174. The recombinant bearing a SHLOX insert in the correct orientation was treated with NcoI to release the 842bp fragment consisting of 39 bp of pET3d and 813bp

SHLOX. Following gel purification, the resulting linear plasmid was recircularised and cloned in *E.coli* HMS174. Clones were sequenced. The verified construct (designated p3DLox) contained the 297bp SHLOX 3'-terminal region encoding a 13,653 Da polypeptide (designated Dlox) corresponding to a carboxyterminal sequence of human lysyl oxidase.

5. p3dL^{m2}

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Sequences of both the $3aL^{ml}$ and the $12aL^{ml}$ recombinant polypeptides were not considered to precisely 10 reflect the primary structure of mature human lysyl oxidase. Each had N-terminal extensions (of the order of 14 residues for $3aL^{ml}$ and 24 residues for $12aL^{ml}$) originating as a consequence of cloning and derived from their vector sequences. These were foreign to mature 15 human lysyl oxidase. To minimise the possible interference of these extensions on the properties and structure of the recombinant mature protein, it was considered worthwhile to place the translational start site as close as possible to aspartate $_{144}$ of SHLOX. 20 Accordingly use was made of an internal methionine 141 residue fortuitously located just three residues upstream of aspartate₁₄₄ comprising the proposed N-terminus of the processed protein. Although there was no convenient recognition site in this region, and in particular no 25 NcoI or NdeI endonuclease restriction site in this part of the SHLOX sequence, an appropriate modification was planned by site-directed mutagenesis. The plasmid chosen for mutagenesis was pGEXSHLOX. pGEXSHLOX is identical to pGEXLOX-2T. An Ncol primer was designed (CCC GTG TTG ACG 30 CCA TGG TTG GTG) which generated two nucleotide changes to transform the GGT ATG GTT target sequence into GCC ATG GTT. After generating the mutation, the resulting construct had two NcoI sites; the second (downstream) site was already in the SHLOX sequence. This offered a 35 simple clone construction scheme. After screening, the mutant clone was used to prepare purified plasmid by

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CsCl-ethidium bromide density gradient centrifugation. From this DNA the 396 bp fragment, flanked by NcoI sites at positions 419 to 424 and 815 to 820 of the mutated SHLOX, was excised. This fragment was then inserted into pDLox. The orientation and primary structure of the recombinant was confirmed by sequencing. construct, designated $p3dL^{m2}$, was electroporated into E.coli BL21/DE3. The inventor recognises that the Asp 144 form can be generated by, for example, treatment of any of the pGEXLOXm1 or p3aLm1 or p12aLm1 forms with an appropriate protease, for example procollagen Cproteinase.

Yeast constructs 6.

Various constructs were made for expression in yeast, including pCNNinv (Carlsberg Research Laboratories) and pYEX-BX (AMRAD). Pichia was chosen as an alternative expression host. For pCNNinv, a form of the synthetic gene corresponding to mature secreted lysyl oxidase was generated by PCR from pl2aLml, using primers which included at their 5' termini EagI sites. 20 . amplification product was purified from an agarose gel using agarase, ligated to pGEM-T and transformed into E. coli XL1-Blue cells. Nine colonies from 16 candidate clones generated DNA that upon cutting with EagI contained bands of approx. 750 and 3004 bp. Two colonies were selected for further investigation. Manual sequencing of one of the two recombinants showed a deletion. The other clone was correct over the sequenced region. A Qiagen (GmbH, Germany) preparation of DNA from this clone was cut with EagI in parallel with pCNNinv. Both forms of DNA were purified using agarase. pCNNinv was treated with calf intestinal alkaline phosphatase and purified by phenol/chloroform extraction. The vector and insert were ligated and transformed into E. coli XL1-Blue cells. Twenty colonies were screened using HindIII, and two colonies appeared to have inserts in the correct orientation. Spheroplasts of

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Saccharomyces cerevisiae strain DBY747 (AMRAD) were transformed separately with the two constructs, and one of these was subsequently chosen for detailed analysis. Complete DNA sequencing confirmed the correct sequence.

For cloning into pYEX-BX, the vector was cleaved with BamHI and dephosphorylated using calf intestinal alkaline phosphatase. Separately, pGEXLOX-2T was treated with restriction enzymes NotI and BamHI and the SHLOX fragment was purified from an agarose gel. This fragment was ligated to the linearised vector, after which ends were filled using deoxynucleoside triphosphates and Sequenase v2.0 (Amersham). Blunt end ligation was then carried out. Correct constructs were identified by restriction enzyme digestion followed by complete sequencing of the fragment and adjacent vector sequences.

All restriction enzymes and agarase used in this study were obtained from Boehringer Mannheim, Germany.

Preparation of various soluble proteins

Growth of E. coli cultures for recombinant lysyl oxidase preparation

A single colony from an agar plate was grown overnight with gyrotary shaking in 3ml LB + ampicillin media, and the entire culture was used to inoculate 250ml 2TY medium containing $70\mu \mathrm{gml}^{-1}$ ampicillin in a 1L conical flask. The inoculated medium was grown at 37°C with vigorous shaking (280-290rpm) until OD $_{600}$ ~0.6-0.8 (about 2h) upon which expression was induced by the addition of IPTG (usually 0.4mM IPTG for DLox, 3aLml and 3dLm2; 0.1mM IPTG for GST-SHLOX). Prior to induction a 0.5ml aliquot of the cell culture was withdrawn for analysis by SDS-PAGE and to confirm the absence of inclusion bodies by phase contrast microscopy. The induced cell culture was incubated under the same conditions for a further 2.5 to 3 hours then transferred to 4°C for 30 minutes before harvesting. At this point a $4\mu l$ aliquot was taken for analysis by phase contrast microscopy for the presence of

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inclusion bodies. If inclusion bodies were present, this was considered a reliable indicator of successful expression. Otherwise expression was monitored by SDS-PAGE of total bacterial protein alongside a sample collected before induction.

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The culture (250 ml, 4°C) was divided into 50 ml aliquots in polypropylene tubes and cells were harvested by centrifugation (10 minutes, 5,000 rpm, 4°C). The supernatant was thoroughly removed and the wet pellets were weighed. A typical yield was 0.5 to 1g packed wet cells per 50 ml culture. Pellets were either stored at -80°C for several months, or used immediately for inclusion body purification.

Purification of inclusion bodies

Thawed pellets were thoroughly resuspended on ice in ~10 vol/weight of lysis buffer (50mM TrisHCL pH8.0, lmM EDTA, 100mM NaCl (usually 5 to 7 ml per tube) and lysed at 4°C with 1mg/ml lysozyme (added in the form of a 10times solution in lysis buffer) for 30 minutes with constant mixing. Mixing was achieved by continuous mechanical inversion on a vertical platform at 30 to 40 To inhibit putative subsequent protease activity PMSF was then added to 0.5 mM. Quantitative cell 10 disruption was achieved by incubation with 1% Triton-X100 (4°C, constant mixing, 30 minutes). Cell lysates showed high viscosity because of released bacterial chromosomal To reduce this effect DNA was digested until visual loss of viscosity with 0.1mg/ml DNase in the presence of 15 $10 \mathrm{mM} \ \mathrm{MgCl}_2$ with constant mixing (typically 20 to 30 minutes at 4°C). Viscosity was further reduced by sonication using three bursts of 10 seconds each in tubes immersed in ice. The extent of lysis was checked by phase contrast microscopy and if necessary the lysis time 20 was extended and/or further sonication applied.

After centrifugation (typically 5,000 rpm, 0° to 4°C, 20 minutes) the supernatant was carefully decanted and residual drops of supernatant were removed with a tissue, taking care not to disturb the pellet. Ice-cold lysis buffer (volume equal to that of the the discarded supernatant) was mixed with the pellet containing inclusion bodies, and the pellet was dispersed either by repeated passage through a blue Gilson Pipetman or by 5 to 6 seconds sonication with a microtip at nearly maximal setting.

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A second centrifugation was similarly performed and the supernatant was discarded. The washed pellet was resuspended in an equal volume of ice-cold 2M urea in 10mM potassium phosphate buffer, pH 8.2, and incubated for 10 minutes with constant mixing at 4°C, followed by centrifugation at 5,000 rpm, 4°C, 20 minutes to collect purified inclusion body fraction in the separated pellet.

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(unless frozen) in the presence of 6M urea at 4°C.

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Solubilisation of inclusion body protein

Solubilisation was achieved immediately after purification using ice-cold U6KP (6M urea in 10 mM potassium phosphate buffer, pH 8.2). The solution was autoclaved and stored at room temperature prior to use. The amount of U6KP used for solubilisation determined the eventual concentration of recombinant lysyl oxidase. Typically 5 ml U6KP was used to solubilise the pellet derived from 50 ml bacterial culture, to yield 2 to 3mg/ml total protein. Mercaptoethanol was added to 2 to 3 mM. To help solubilisation mechanical disruption with a plastic Gilson tip was initially tried but this often led to loss of material which adhered to the inside and outside of the tip. Instead, the mixture was usually mechanically mixed by continuous mechanical inversion on a vertical platform at 30 to 40 rpm overnight, or until there was no sign of clumps. At all times the temperature was kept ~0° to 4°C.

The total protein concentration was estimated by Bradford assay, and CuCl₂ was added at ⁵ to 10 times molar excess over estimated protein. This was usually 200 µM CuCl₂. Solubilised protein stocks were used directly for protein analysis or for refolding of the protein by dialysis in various buffers, including phosphate buffers, depending on the nature of the experiment.

Refolding of inclusion bodies by dialysis against phosphate buffer

For dialysis against phosphate buffer 10 mm and 32 mm dialysis tubing (Selby, Australia) was usually used. Cut tubing was boiled in MilliQ water for 10 minutes and

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briefly soaked in dialysis buffer prior to use. 10mM or 16 mM phosphate buffer was prepared from 1M $\rm KH_2PO_4$ and 1M $\rm K_2HPO_4$ solution stocks and chemical grade distilled water, with the occasionally pH adjusted with 1M KOH.

Dialysis of 2 or 3 ml of a 6M urea solution of recombinant protein (from 2 or 3 tubes, respectively) was routinely performed against 2 L urea-free buffer, with buffer changes after each dialysis. Copper-containing samples were first dialysed against buffer containing an approximately equimolar concentration of CuCl_2 (usually $10\text{--}50~\mu\text{M}$), in the expectation that removal of urea would be accompanied by integration of copper and bias towards a conformationally relevant form of the protein. Excess of copper was then removed in the following dialysis.

After two consecutive dialyses (one against a copper-containing buffer, and the other against a copper-free buffer at pH 9.6) the dialysed material was transferred into polypropylene tubes on ice. Soluble and insoluble fractions were separated by centrifugation and analysed by SDS-PAGE. The pellet and the supernatant obtained after dialysis was stored at 4°C without obvious protein degradation for at least 3 weeks, but some solubility changes were noticed during storage.

Protein from S. cerevisiae cultures

To obtain lysyl oxidase in yeast cell pellets and lysates, a colony was grown in CSM-leu for 48 hours. To purify secreted protein, cells were separated from growth media by centrifugation, and the supernatant was dialysed against ice-cold U6KP. Cells were washed in 5mM KPB (retained) then resuspended in 5mM KPB plus lyticase. After incubation for 1 hour at 30°C, cells were sonicated until apparent lysis as detected by phase contrast microscopy. The pellet and lysate were separated by centrifugation. The pellet and lysates were washed with U6KP and dialysed separately against ice-cold U6KP then against KPB, pH 9.6.

280nm.

PROTEIN STUDIES

To confirm the identity of the expressed protein, in each case DNA sequences were determined. Purified soluble recombinant lysyl oxidase forms were all examined by

- SDS-PAGE, electrospray mass spectrometry, amino acid analysis and N-terminal protein sequencing to confirm the identity of the overexpressed protein. Detailed analyses of full-length protein demonstrated the following:
- (1) the protein bound copper at ~ 1 mole Cu²⁺ per mole 10 protein; this assay was performed by atomic absorption spectroscopy on a Varian SpectrAA-10. Protein samples were examined for protein content using a Bradford assay at 595 nm, and the ratio molar content of copper to protein was determined.
- 15 (2) the protein exists as multimers in solution; multimer formation was observed by column chromatography in experiments using either Sephadex G100 or G200, compared with protein standards. Aliquots were examined spectrophotometrically at 280nm. Light scattering
- studies (Malvern Instruments S4700 version PCS v 1.26) showed size distributions, including approximately spherical particles at 30 to 56 nm.
 - (3) multiple peak formation by ion-exchange chromatography; multiple peaks were observed for 3dLm2 when analysed on a 100 nm x 4.6 mm HperD Q column on a BioSys2000 system (Beckman, USA), using a linear NaCl gradient from) to 0.4 M NaCl, over 9 min in the presence of 6M urea and 16 mM potassium phoshpate buffer, at
- 30 (4) organic cofactor formation as evidenced by redox-cycling (Paz et al., 1991), covalent radiolabel attachment of 14-C-ethylenediamine, and Raman spectroscopy of the p-nitrophenylhydrazone
- (5) enzyme activity including conversion of recombinant35 tropoelastin to aqueous-insoluble material.

In the yeast expression product studied characteristics 2 and 5 have been verified indicating that functional protein is being produced while all 5

characteristics have been verified for E.coli expression particularly with respect to the p3dL m2 construct.

Levels of expression obtained have been approximately 30% of total cell protein in $E.\ coli$ and 10% of total cell protein in yeast.

Benzylamine substrate assay

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The assay was performed according to the method of Trackman et al. 1979. The assay monitors at 250nm the production of benzylaldehyde from benzylamine in potassium phosphate buffer at 37°C. Lysyl oxidase activity is measured spectrophotometrically at 37°C by the conversion of benzylamine to benzaldehyde at 250nm.

Three samples containing lysyl oxidase and benzylamine showed a slight increase in absorbance at 250nm. In this assay the lysyl oxidase was derived from the plasmid p3dL^{m2}. Controls, including boiled lysyl oxidase and benzylamine, benzylamine and dialysis buffer (no enzyme) and lysyl oxidase and KPB (no substrate) showed no increase in absorbance at 250nm.

Averaging the change in absorbance at 250nm of the three lysyl oxidase and benzylamine samples and estimating the concentration of lysyl oxidase in each assay the specific activity of the enzyme was calculated at 0.0145 umol/min/mg. Wang et al. reports a specific activity of 0.019 U/mg.

Tropoelastin substrate assay

Tropoelastin resuspended in either KPB (10mM,pH8) or PBS (10mM PB 150mM NaCl) was mixed at a final concentration of 10mg/mL, with lysyl oxidase and placed at 37°C for 2 hrs. In this assay, the lysyl oxidase was derived from the p3dL^{m2} plasmid. Controls included tropoelastin + lysyl oxidase + BAPN, tropoelastin + boiled lysyl oxidase, tropoelastion + dialysis buffer, tropoelastin + buffer.

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Equal amounts from each reaction were run on an 8% On staining with Coomassie less intense tropoelastin bands were seen in the tropoelastin in KPB + LO and tropoelastin in KPB + lysyl oxidase + BAPN sample lanes compared to other lanes. This may be indicative of the presence of crosslinked tropoelastin which is too large to enter the gel. Whilst BAPN should inhibit lysyl oxidase activity, preincubation of the enzyme with BAPN for at least 2 hours at 37°C may be required prior to assaying in order for the inhibition to be effective.

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The above 2 samples and the tropoelastin + buffer control were centrifuged, the supernatant removed and the precipitate sent for amino acid analysis. The 2 samples showed a decrease in percentage of total lysine compared to the control. Tropoelastin + buffer contained 6% lysine, tropoelastin + lysyl oxidase 2.5% lysine and tropoelastin + lysyl oxidase + BAPN 2.2% lysine. consistent with the removal of lysine residues to form cross-links such as desmosines. Bedell- Hogan et al reports the presence of 5.6% lysine residues in recombinant tropoelastin compared to 2.4% in alkali insoluble recombinant tropoelastin.

In summary, the results show a lowering of the percentage of lysine in tropoelastin, as determined by the amino acid analysis. Furthermore, an insoluble protein was produced, consistent with the production of an elastin or elastin-like product.

Tritiated tropoelastin assay

Recombinant tropoelastin was prepared essentially as described Martin et al. 1995, except that BL21 cells containing pSHELF (WO 94/14958) were grown in "complete synthetic medium without lysine" (CSM-Lys), and supplemented with L-[4,5-tritium] lysine monochloride (Amersham) at 83 Ci/mmol. Accordingly, the resulting tropoelastin was radiolabelled to approximately 40,000,000 cpm/mg protein. This was lyophilised, then redissolved in assay buffer (0.1M Na₂B₄O₇, 0.15M NaCl,

 $\mathtt{pH8})$ and centrifuged through Millipore Ultrafree MC for 1 hour at 7500 rpm at 4°C. The retentate was used for subsequent assays. Assays were then performed by incubation with lysyl oxidase, and assessed according to 5 the protocol of Shackelton and Hulmes, 1990. assay, the lysyl oxidase was derived from the $p3dL^{m2}$ plasmid. Tritiated tropoelastin was mixed with lysyl oxidase and left at 37°C for 16 hours. Tritiated water formed during the incubation, an indication of lysyl oxidase activity, was isolated by ultrafiltration and 10 radioactivity quantified by liquid scintillation spectrometry. A control for endogenous lysyl oxidaselike activity in E. coli was generated by taking an E. coli strain containing the pET3d plasmid through the same purification procedure as that used to produce lysyl 15 oxidase. A sample of this pET3d control was also mixed with tritiated tropoelastin. Preliminary results are consistent with some lysyl oxidase-like activity. cases samples containing lysyl oxidase produced more 20 tritiated water than those containing pET3d (Figure 5).

INDUSTRIAL APPLICABILITY

The genetic constructs of the present invention are useful in the preparation of high yields of biocompatible biospecific lysyl oxidase useful in vivo applications and 25 other situations where catalysis of amine oxidation is required. The expression products of the invention are useful as agents for crosslinking of (but not confined to) tropoelastin and collagen both in vitro and in vivo, to join primary-amine materials e.g. in surgical or veterinary applications including natural and artificial 30 materials, regulating cell growth, interfering with/promoting fibrosis, oxidising accessible lysines (and other amines) to modify behaviour of those molecular components and as components of assay systems designed to detect/quantify amines. Further, the genetic constructs 35 and expression products of the invention have potential in cancer therapy.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: The University of Sydney, N/A N/A
 - (ii) TITLE OF INVENTION: SYNTHETIC POLYNUCLEOTIDES
 - (iii) NUMBER OF SEQUENCES: 30
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: GRIFFITH HACK
 - (B) STREET: LEVEL 8, 168 WALKER STREET
 - (C) CITY: NORTH SYDNEY
 - (D) STATE: NEW SOUTH WALES
 - (E) COUNTRY: AUSTRALIA
 - (F) ZIP: 2060
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: AU PO1565
 - (B) FILING DATE: 09-AUG-1996
 - (C) CLASSIFICATION:
 - (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: AU PO1565
 - (B) FILING DATE: 09-AUG-1996
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Kurts, ANN D
 - (B) REGISTRATION NUMBER: N/A
 - (C) REFERENCE/DOCKET NUMBER: 24078DM:ADK
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 61 2 9957 5944
 - (B) TELEFAX: 61 2 9957 6288
 - (C) TELEX: AA 26547
 - (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1251 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "nucleic acid"
 - (iii) HYPOTHETICAL: YES
 - (iv) ANTI-SENSE: YES
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATGCGCTTCG CCTGGACCGT GCTCCTGCTC GGGCCTTTGC AGCTCTGCGC GCTAGTGCAC

TGCGCCCCTC CCGCCGCGG CCAACAGCAG CCCCCGCGCG AGCCGCCGGC GGCTCCGGGC

GCCTGGCGCC AGCAGATCCA ATGGGAGAAC AACGGGCAGG TGTTCAGCTT GCTGAGCCTG	180
GGCTCACAGT ACCAGCCTCA GCGCCGCCGG GACCCGGGCG CCGCCGTCCC TGGTGCAGCC	240
AACGCCTCCG CCCAGCAGCC CCGCACTCCG ATCCTGCTGA TCCGCGACAA CCGCACCGCC	300
GCGGCGCGAA CGCGGACGGC CGGCTCATCT GGAGTCACCG CTGGCCGCCC CAGGCCCACC	360
GCCCGTCACT GGTTCCAAGC TGGCTACTCG ACATCTAGAG CCCGCGAACG TGGCGCCTCG	420
CGCGCGGAGA ACCAGACAGC GCCGGGAGAA GTTCCTGCGC TCAGTAACCT GCGGCCGCCC	480
AGCCGCGTGG ACGGCATGGT GGGCGACGAC CCTTACAACC CCTACAAGTA CTCTGACGAC	540
AACCCTTATT ACAACTACTA CGATACTTAT GAAAGGCCCA GACCTGGGGG CAGGTACCGG	600
CCCGGATACG GCACTGGCTA CTTCCAGTAC GGTCTCCCAG ACCTGGTGGC CGACCCCTAC	660
TACATCCAGG CGTCCACGTA CGTGCAGAAG ATGTCCATGT ACAACCTGAG ATGCGCGGCG	720
GAGGAAAACT GTCTGGCCAG TACAGCATAC AGGGCAGATG TCAGAGATTA TGATCACAGG	780
GTGCTGCTCA GATTTCCCCA AAGAGTGAAA AACCAAGGGA CATCAGATTT CTTACCCAGC	840
CGACCAAGAT ATTCCTGGGA ATGGCACAGT TGTCATCAAC ATTACCACAG TATGGATGAG	900
TTTAGCCACT ATGACCTGCT TGATGCCAAC ACCCAGAGGA GAGTGGCTGA AGGCCACAAA	960
GCAAGTTTCT GTCTTGAAGA CACATCCTGT GACTATGGCT ACCACAGGCG ATTTGCATGT	1020
ACTGCACACA CACAGGGATT GAGTCCTGGC TGTTATGATA CCTATGGTGC AGACATAGAC	1080
TGCCAGTGGA TTGATATTAC AGATGTAAAA CCTGGAAACT ATATCCTAAA GGTCAGTGTA	1140
AACCCCAGCT ACCTGGTTCC TGAATCTGAC TATACCAACA ATGTTGTGCG CTGTGACATT	1200
CGCTACACAG GACATCATGC GTATGCCTCA GGCTGCACAA TTTCACCGTA T	1251
(2) INFORMATION FOR SEC. TO US	

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 417 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: YES
- (iv) ANTI-SENSE: NO
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
- Met Arg Phe Ala Trp Thr Val Leu Leu Leu Gly Pro Leu Gln Leu Cys
 1 10 15
- Ala Leu Val His Cys Ala Pro Pro Ala Ala Gly Gln Gln Pro Pro
 20 25 30
- Arg Glu Pro Pro Ala Ala Pro Gly Ala Trp Arg Gln Gln Ile Gln Trp
- Glu Asn Asn Gly Gln Val Phe Ser Leu Leu Ser Leu Gly Ser Gln Tyr 50 60

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Gln 65	Pro	Glı	ı A	rg	Arg	70	g A	sp :	Pro	Gl	у 1	Ala	Ala 75	۷a	.1 I	?ro	G13	7 #	Ma.	80 818	1
Asn	Ala	Se	r A	la	Glr 85	G1	n P	ro.	Arg	Th	r !	Pro 90	Ile	Le	eu I	Leu	Ile	e 1	Arg 95	Ası	þ
Asn	Arg	Th		la 00	Ala	LA 1	a A	rg	Thr	Ar 10	g ')5	Thr	Ala	G3	ly :	Ser	Se:	r (Va	
Thr	Ala	G1 11		Arg	Pro) Ai	g P	ro	Thr 120	A)	la.	Arg	His	T	ср	Phe 125	G1	n i			
Tyr	Ser 130		r s	Ser	Ar	A E	la A	rg .35	Glu	A	rg	Gly	Ala	1 Se	er 40	Arg	Al	a	Glu	Аs	n
Gln 145	Thr	Al	.a 1	Pro	Gl	y G:	lu \ 50	/al	Pro	A.	la	Leu	Ser 155	: A	sn	Leu	Ar	g	Pro	Pr 16	0
Ser	Arg	y Va	al.	qaA	Gl 16	у М 5	et '	Val	GJÀ	, A	qa	Asp 170	Pro	т	уr	Asn	Pı	0	Tyr 175	L	/S
Tyr	Se	c As	sp	Asp 180	As	n P	ro'	Tyr	Тут	r A 1	sn 85	Tyr	Ту	r A	qa	Thr	15	/r 90	Glu	Aı	rg
Pro	Ar	g P:	ro 95	Gly	G1	уА	rg '	Tyr -	Arg 200	g P O	ro	Gly	ту	r G	ly	Thr 205	G:	ly	Tyr	Pl	ne
Glr	Ty 21		ly	Leu	Pi	:0 A	ap	Leu 215	۷a	1 <i>P</i>	la	Asp	Pr	0 7	Гуг 220	Ту	: I	le	Gln	A	la
Ser 229	Th	r T	yr	Val	. G:	n I	Lys 230	Met	Se	r N	1et	Туз	23	n I	Leu	Arg	g C	уs	Ala	. A 2	1a 40
Glu	ı Gl	u A	sn	Cys	3 Le	eu 1 15	Ala	Ser	Th	rl	Ala	Ty:	r Ar	g)	Ala	As	pV	al	Arg 255	A 5	ga.
Ty	r As	рН	: (is	Arg 26	g V.	al :	Leu	Leu	ı Ar	g i	Phe 265	Pro	o Gl	n	Arg	Va	1 L 2	ys !70	Ası	n G	ln
Gl.	y Tì	ır S	er 175	As	p P	he	Leu	Pro	Se 28	er .	Arg	, Pr	o Ai	rg '	Туг	Se 28	r 1 5	rp	Gl	u I	rp
Hi		er (Сув	Hi	s G	ln	His	Ту: 29!	r Hi 5	is	Sei	. Me	t A	вр	Gl: 300	ı Ph	e S	Ser	Hi.	s 7	ſyr
As 30		eu J	Leu	As	pА	la	Asn 310	Th	r G	ln	Ar	g Ar	g V	al 15	Ala	a Gl	.u (313	/ Hi	s :	Lys 320
Al	a S	er 1	Phe	: Су	s I	eu 25	Glu	As	p Tl	hr	Se	r Cy 33	rs A 80	sp	Ty:	r Gl	ly '	Ту	r Hi 33	.s . 15	Arg
Aı	g P	he .	Ala	2 Cy 34	rs 7	hr	Ala	Hi	s T	hr	G1 34	n G1 5	ly L	eu	Se	r Pi	ro	G1; 35	у С <u>У</u>	rs	Tyr
As	sp T		Ту: 35!		ly i	Ala	Авр	11	.е А 3	qa. 00	Су	s G	ln I	,rb	Il	e A	вр 65	Il	e Tì	ır	qaA
V		ys 70	Pro	o G:	ly i	Asn	Туз	37	.e L 75	eu	Ly	s V	al S	Ser	Va 38	1 A	sn	Pr	o Se	er	Туг
	eu 1 85	al	Pr	o G	lu	Ser	Aei 390	, Σ	r I	hr	Αε	sn A	en 7	/al 395	Va	l A	rg	Су	s A	sp	11e 400
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(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1184 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "nucleic acid"
- (iii) HYPOTHETICAL: YES
- (iv) ANTI-SENSE: YES
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

2 == 0.5151			
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TCCAGTGGGA AAACAACGGT CAGGTTTTCT CCCTGCTGTC C	CCTGGGTTCC	CAGTACCAGC	120
CGCAGCGTCG TCGTGACCCG GGTGCTGCTG TTCCGGGTGC T	TGCTAACGCT	TCCGCTCAGC	180
AGCCGCGTAC CCCGATCCTG CTGATCCGTG ACAACCGTAC C	CGCGGCCGCT	CGTACCCGTA	240
CCGCTGGTTC CTCCGGTGTT ACTGCAGGTC GTCCGCGTCC G	GACCGCGCGC	CACTGGTTCC	300
AGGCTGGTTA CTCCACCTCC CGTGCTCGTG AAGCTGGTGC T	TCCCGTGCT	GAAAACCAGA	360
CCGCTCCGGG TGAAGTTCCA GCGCTGTCCA ACCTGCGTCC G	GCCGTCCCGT	GTTGACGGTA	420
TGGTTGGTGA CGACCCGTAC AACCCGTACA AGTACTCCGA C	GACAACCCG	TACTACAACT	480
ACTACGACAC CTACGAGCGC CCGCGTCCGG GTGGTCGTTA C	CCGTCCGGGT	TACGGTACCG	540
GTTACTTCCA GTACGGTCTG CCGGACCTGG TTGCTGACCC G	TACTACATC	CAGGCTTCCA	600
CCTACGTTCA GAAAATGTCC ATGTACAACC TGCGTTGCGC TG	GCTGAAGAA .	AACTGCCTGG	660
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CGCAGCGTGT TAAAAACCAG GGCACCTCCG ACTTCCTGCC G	TCCCGTCCG	CGTTACTCCT	780
GGGAATGGCA CTCCTGCCAC CAGCACTACC ACTCCATGGA CO	GAATTCTCC (CACTACGACC	840
TGCTGGACGC TAACACCCAG CGTCGTGTTG CTGAAGGTCA C	AAAGCTTCC :	TTCTGCCTGG	9 0 0
AAGACACCTC CTGCGACTAC GGTTACCACC GTCGTTTCGC TT	TGCACCGCT (CACACCCAGG	960
GTCTGTCCCC GGGTTGCTAC GACACCTACG GTGCTGACAT CO	GACTGCCAG 1	rggatcgaca	1020
TCACCGACGT TAAACCGGGT AACTACATCC TGAAAGTTTC CG			1080
TTCCGGAATC CGACTACACC AACAACGTGG TTCGTTGCGA TA			1140
ACGCTTACGC TTCCGGTTGC ACCATCTCCC CGTACTAATG AT			1184
/3\ TX700x40=			

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 391 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(iii) HYPOTHETICAL: YES

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Ser Gln Gln Gln Pro Pro Arg Glu Pro Pro Ala Ala Pro Gly Ala Trp

Arg Gln Gln Ile Gln Trp Glu Asn Asn Gly Gln Val Phe Ser Leu Leu 20 25 30

Ser Leu Gly Ser Gln Tyr Gln Pro Gln Arg Arg Arg Asp Pro Gly Ala

Ala Val Pro Gly Ala Ala Asn Ala Ser Ala Gln Gln Pro Arg Thr Pro 50 55

Ile Leu Leu Ile Arg Asp Asn Arg Thr Ala Ala Ala Arg Thr Arg Thr 65 70 75 80

Ala Gly Ser Ser Gly Val Thr Ala Gly Arg Pro Arg Pro Thr Ala Arg 85 90 95

His Trp Phe Gln Ala Gly Tyr Ser Thr Ser Arg Ala Arg Glu Ala Gly 100 105

Ala Ser Arg Ala Glu Asn Gln Thr Ala Pro Gly Glu Val Pro Ala Leu 115 120 125

Ser Asn Leu Arg Pro Pro Ser Arg Val Asp Gly Met Val Gly Asp Asp 130 135

Pro Tyr Asn Pro Tyr Lys Tyr Ser Asp Asn Pro Tyr Tyr Asn Tyr
145 150 155 160

Tyr Asp Thr Tyr Glu Arg Pro Arg Pro Gly Gly Arg Tyr Arg Pro Gly
165 170 175

Tyr Gly Thr Gly Tyr Phe Gln Tyr Gly Leu Pro Asp Leu Val Ala Asp

Pro Tyr Tyr Ile Gln Ala Ser Thr Tyr Val Gln Lys Met Ser Met Tyr 195 200 205

Asn Leu Arg Cys Ala Ala Glu Glu Asn Cys Leu Ala Ser Thr Ala Tyr 210 215 220

Arg Ala Asp Val Arg Asp Tyr Asp His Arg Val Leu Leu Arg Phe Pro

Gln Arg Val Lys Asn Gln Gly Thr Ser Asp Phe Leu Pro Ser Arg Pro

Arg Tyr Ser Trp Glu Trp His Ser Cys His Gln His Tyr His Ser Met

Asp Glu Phe Ser His Tyr Asp Leu Leu Asp Ala Asn Thr Gln Arg Arg 285

Val Ala Glu Gly His Lys Ala Ser Phe Cys Leu Glu Asp Thr Ser Cys 290 295 300

Asp Tyr Gly Tyr His Arg Arg Phe Ala Cys Thr Ala His Thr Gln Gly 305 310 320

---- (TIU F 06)

I	eu	Ser	Pro	Gly	Cys 325	Tyr	Asp	Thr	Tyr	Gly 330	Ala	Asp	Ile	Asp	Cys 335	Gln	
T	rp	Ile	Asp	Ile 340	Thr	Asp	Val	Lys	Pro 345	Gly	Asn	Tyr	Ile	Leu 350	Lys	Val	
S	Ger	Val	Asn 355	Pro	Ser	Tyr	Leu	Val 360	Pro	Glu	Ser	Asp	Tyr 365	Thr	Asn	Asn	
V	al	Val 370	Arg	Сув	Asp	Ile	Arg 375	Tyr	Thr	Gly	His	His 380	Ala	Tyr	Ala	Ser	
	1y 85	Cys	Thr	Ile	Ser	Pro 390	Tyr										
(2) IN	FOR	MATI	ON I	FOR S	EQ I	D NC):5:										
(i)	(A) (B) (C)	LEN TYI STI	CHA IGTH: PE: r PANDE POLOG	95 ucle DNES	base ic a S: s	pai cid ingl	rs									
(i	i) :	MOLE (A)	CULE	TYP	E: c	ther : /d	nuc lesc	leic = "s	aci	d etic	oli	.gonu	ıcleo	tide	. "		
(ii	i) :	HYPC	THET	'ICAL	: YE	S		_									
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(x	i) :	SEQU	ENCE	DES	CRIP	TION	: SE	Q ID	NO:	5 :	,						
GATCCC	AGC	A GC	AGCC	GCCG	CGT	GAAC	CGC	CGGC	TGCT	CC G	GGTG	CTTG	G CG	TCAG	CAGA		60
TCCAGT	GGG	AA A	ACAA	CGGT	CAG	GTTT	TCT	CCCT	G								95
(2) IN	FORI	MATI	ON F	OR S	EQ I	D NO	:6:										
(:	i) \$	(A) (B) (C)	LEN TYP STR	CHA GTH: E: n ANDE OLOG	85 ucle DNES	base ic a S: s	pai cid ingl	rs									
(ii	i) N	MOLE (A)	CULE DES	TYP: CRIP	E: o	ther :/d	nuc esc	leic = "s	aci ynth	d etic	oli	gonu	cleo	tiđe	n		
(iii	i) F	IYPO	THET	ICAL	: YE	s											
(iv	r) I	MTI	- SEN	SE:	YES												
(xi	i) s	EQUI	ENCE	DES	CRIP	rion	: SE	Q ID	NO:	6:				•			
CTGTCCC	TGG	GT	rccci	AGTA	CCA	GCCG(CAG (CGTC	GTCG'	rg a	CCCG	GGTG	C TG	CTGT	rccg		60
GGTGCTG																	85
(2) INF	ORM	TATIO	ON FO	OR SI	EQ II	NO:	:7:										
(i	.) S	(A) (B)	LENG	CHAI FTH: E: nu ANDEI	86 h clei	ase C a	pair	rs						-			

(D) TOPOLOGY: linear	-
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
AGCCGCGTAC CCCGATCCTG CTGATCCGTG ACAACCGTAC CGCGGCCGCT CGTACCCGTA	60
CCGCTGGTTC CTCCGGTGTT ACTGCA	86
(2) INFORMATION FOR SEQ ID NO:8:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 93 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	
GGTCGTCCGC GTCCGACCGC GCGCCACTGG TTCCAGGCTG GTTACTCCAC CTCCCGTGCT	
CGTGAAGCTG GTGCTTCCCG TGCTGAAAAC CAG	93
(2) INFORMATION FOR SEQ ID NO:9:	
(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 91 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	-
ACCGCTCCGG GTGAAGTTCC AGCGCTGTCC AACCTGCGTC CGCCGTCCCG TGTTGACGGT	6
ATGGTTGGTG ACGACCCGTA CAACCCGTAC A	9
(2) INFORMATION FOR SEQ ID NO:10:	
(i) SEQUENCE CHARACTERISTICS:	

11 0 20/00030

(A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:	
AGTACTCCGA CGACAACCCG TACTACAACT ACTACGACAC CTACGAGCGC CCGCGTCCGG	60
GTGGTCGTTA CCGTCCGGGT TACGGTAC	88
(2) INFORMATION FOR SEQ ID NO:11:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	,
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:	
CGGTTACTTC CAGTACGGTC TGCCGGACCT GGTTGCTGAC CCGTACTACA TCCAGGCTTC	60
CACCTACGTT CAGAAAATGT CCATGTAC	88
(2) INFORMATION FOR SEQ ID NO:12:	0.6
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 97 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:	
ACCTGCGTT GCGCTGCTGA AGAAAACTGC CTGGCTTCCA CCGCTTACCG TGCTGACGTT	60
CGTGACTACG ACCACCGTGT TCTGCTGCGT TTCCCGC	97

(2) INFORMATION FOR SEQ ID NO:13:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 99 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
AGCGTGTTAA AAACCAGGGC ACCTCCGACT TCCTGCCGTC CCGTCCGCGT TACTCCTGGG	60
AATGGCACTC CTGCCACCAG CACTACCACT CCATGGACG	99
(2) INFORMATION FOR SEQ ID NO:14:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 99 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid(A) DESCRIPTION: /desc = "synthetic oligonucleotide"	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	_0
AATTCTCCCA CTACGACCTG CTGGACGCTA ACACCCAGCG TCGTGTTGCT GAAGGTCACA	60
AAGCTTCCTT CTGCCTGGAA GACACCTCCT GCGACTACG	99
(2) INFORMATION FOR SEQ ID NO:15:	
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<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	

GTTACCACCG TCGTTTCGCT TGCACCGCTC ACACCCAGGG TCTGTCCCCG GGTTGCTACG

ACACCTACGG TGCTGACATC GACTGCCAGT GGA	93
(2) INFORMATION FOR SEQ ID NO:16:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 93 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SECHENCE DESCRIPTION CON CONTRACT	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
TCGACATCAC CGACGTTAAA CCGGGTAACT ACATCCTGAA AGTTTCCGTT AACCCGTCCT	60
ACCTGGTTCC GGAATCCGAC TACACCAACA ACG	93
(2) INFORMATION FOR SEQ ID NO:17:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 77 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
TGGTTCGTTG CGATATCCGT TACACCGGTC ACCACGCTTA CGCTTCCGGT TGCACCATCT CCCCGTACTA ATGATAG	60
	77
(2) INFORMATION FOR SEQ ID NO:18:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 85 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:	

BAAAACCTGA CCGTTGTTTI CCCACTGGAT CTGCTGACGC CAAGCACCCG GAGCAGCCGG	60
CGGTTCACGC GGCGGCTGCT GCTGG	85
(2) INFORMATION FOR SEQ ID NO:19:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 85 base pairs (B) Type: nucleic acid	
(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:	60
CGGAAGCGTT AGCAGCACCC GGAACAGCAG CACCCGGGTC ACGACGACGC TGCGGCTGGT	85
ACTGGGAACC CAGGGACAGC AGGGA	85
(2) INFORMATION FOR SEQ ID NO:20:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:	60
GTAACACCGG AGGAACCAGC GGTACGGGTA CGAGCGGCCG CGGTACGGTT GTCACGGATC	60
AGCAGGATCG GGGTACGCGG CTGCTGAG	88
(2) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 91 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
TTCAGCACGG GAAGCACCAG CTTCACGAGC ACGGGAGGTG GAGTAACCAG CCTGGAACCA	60
GTGGCGCGC GTCGGACGCC GACGACCTGC A	91
(2) INFORMATION FOR SEQ ID NO:22:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 90 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:	
GTTGTACGGG TCGTCACCAA CCATACCGTC AACACGGGAC GGCGGACGCA GGTTGGACAG	60
CGCTGGAACT TCACCCGGAG CGGTCTGGTT	90
(2) INFORMATION FOR SEQ ID NO:23:	
 (A) LENGTH: 91 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:	
CGTAACCCGG ACGGTAACGA CCACCCGGAC GCGGGCGCTC GTAGGTGTCG TAGTAGTTGT	60
AGTACGGGTT GTCGTCGGAG TACTTGTACG G	91
(2) INFORMATION FOR SEQ ID NO:24:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 98 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
GCAGGTTGTA CATGGACATT TTCTGAACGT AGGTGGAAGC CTGGATGTAG TACGGGTCAG	60
CAACCAGGTC CGGCAGACCG TACTGGAAGT ACCGGTAC	98
(2) INFORMATION FOR SEQ ID NO:25:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 97 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: other nucleic acid(A) DESCRIPTION: /desc = "synthetic oligonucleotide"	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
ACACGCTGCG GGAAACGCAG CAGAACACGG TGGTCGTAGT CACGAACGTC AGCACGGTAA	- 60
GCGGTGGAAG CCAGGCAGTT TTCTTCAGCA GCGCAAC	97
(2) INFORMATION FOR SEQ ID NO:26:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	60
AATTCGTCCA TGGAGTGGTA GTGCTGGTGG CAGGAGTGCC ATTCCCAGGA GTAACGCGGA	96
CGGGACGCA GGAAGTCGGA GGTGCCCTGG TTTTTA	,
(2) INFORMATION FOR SEQ ID NO:27:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 88 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
(iii) HYPOTHETICAL: YES	

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(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
GCAGGAGGTG TCTTCCAGGC AGAAGGAAGC TTTGTGACCT TCAGCAACAC GACGCTGGGT	60
GTTAGCGTCC AGCAGGTCGT AGTGGGAG	88
(2) INFORMATION FOR SEQ ID NO:28:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 94 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
GGCAGTCGAT GTCAGCACCG TAGGTGTCGT AGCAACCCGG GGACAGACCC TGGGTGTGAG	60
CGGTGCAAGC GAAACGACGG TGGTAACCGT AGTC	94
(2) INFORMATION FOR SEQ ID NO:29:	24
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 93 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	
(iii) HYPOTHETICAL: YES	
(iv) ANTI-SENSE: YES	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	
TGGTGTAGTC GGATTCCGGA ACCAGGTAGG ACGGGTTAAC GGAAACTTTC AGGATGTAGT	60
FACCCGGTTT AACGTCGGTG ATGTCGATCC ACT	93
(2) INFORMATION FOR SEQ ID NO:30:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 87 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "synthetic oligonucleotide"</pre>	·
(iii) HYPOTHETICAL: YES	

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
GATCCTATCA TTAGTACGGG GAGATGGTGC AACCGGAAGC GTAAGCGTGG TGACCGGTGT	60
AACGGATATC GCAACGAACC ACGTTGT	87

THE CLAIMS

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- 1. A synthetic polynucleotide encoding an amino acid sequence selected from the group consisting of lysyl oxidase, lysyl oxidase-like proteins and variants thereof.
- 2. A synthetic polynucleotide according to claim 1 wherein the synthetic polynucleotide comprises a nucleotide sequence which is derived from a nucleotide sequence expressed in a vertebrate species including a mammalian or avian species.
- 3. A synthetic polynucleotide according to claim 2 wherein the synthetic polynucleotide sequence comprises a nucleotide sequence which is derived from a nucleotide sequence expressed in humans.
- 4. A synthetic polynucleotide according to any one of claims 1-3 wherein the synthetic polynucleotide has a CAI score of at least 0.30 in a particular host cell.
- 5. A synthetic polynucleotide according to claim 4 wherein the particular host cell is selected from the group consisting of *E. coli*, *Saccharomyces cerevisiae* and *Pichia*.
 - 6. A synthetic polynucleotide comprising the nucleotide sequence shown in Figure 2 (SEQ ID NO: 3) or a fragment thereof, wherein the fragment thereof encodes an amino acid sequence which retains the ability of lysyl oxidase or lysyl oxidase-like proteins to catalyse the oxidation of primary amines.
- 7. A synthetic polynucleotide according to claim 6 wherein the nucleotide sequence commences at the codon encoding amino acid residue 141.

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- 8. A synthetic polynucleotide according to claim 6 wherein the nucleotide sequence commences at the codon encoding amino acid residue 144.
- 9. A synthetic polynucleotide according to any one of claims 6 to 8 wherein at least 50% of the nucleotide mutations indicated in Figure 2 (SEQ ID NO:3) have been introduced.
- of claims 6 to 8 wherein the number of nucleotide

 mutations which have been introduced produce a CAI score
 for the synthetic polynucleotide in E. coli of at least
 0.30.
- 11. A synthetic polynucleotide according to any one of claims 1-6 wherein at least 50% of codons for any particular amino acid are selected and altered to reflect preferred codon usage in the particular host.
 - 12. A synthetic polynucleotide according to any one of claims 1-6 wherein at least 50% of codons for amino acids for which altered codons are desirable are altered.
- 20 13. A synthetic polynucleotide according to claim 12 wherein at least 60% of codons for amino acids for which altered codons are desirable are altered.
- 14. A synthetic polynucleotide according to claims
 12 wherein at least 70% of codons for amino acids for
 25 which altered codons are desirable are altered.
 - 15. A synthetic polynucleotide comprising an nucleotide sequence encoding a lysyl oxidase, lysyl oxidase like protein or variant thereof, wherein all or part of the 5' or 3' untranslated region, and/or intronic region of the synthetic polynucleotide is deleted.

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- 16. A synthetic polynucleotide sequence encoding a lysyl oxidase, lsysyl oxidase-like protein or variant thereof, wherein all or part of the region encoding the signal peptide, and/or region encoding the pro-peptide, of the polynucleotide is deleted.
- 17. A synthetic polynucleotide sequence encoding a lysyl oxidase, lsysyl oxidase-like protein or variant thereof, wherein all or part of the region encoding the signal peptide, and/or region encoding the pro-peptide, of the polynucleotide is deleted and wherein all or part of the 5' or 3' untranslated region, and/or intronic region of the synthetic polynucleotide is deleted.
- 18. A synthetic polynucleotide comprising a nucleotide sequence encoding an amino acid sequence selected from the group consisting of lysyl oxidase, lysyl oxidase-like proteins and variants thereof and a further nucleotide sequence wherein the nucleotide sequence encoding lysyl oxidase, lysyl oxidase-like protein or a variant thereof and the further nucleotide sequence are chemically linked so as to encode a single open reading frame.
- 19. A synthetic polynucleotide according to claim 18 wherein the further nucleotide sequence is chemically linked to the 5' end of the nucleotide sequence encoding lysyl oxidase, lysyl oxidase-like protein or a variant thereof.
- 20. A synthetic polynucleotide according to claim 18 or 14 wherein the further nucleotide sequence encodes all or part of a protein which causes the lysyl oxidase,

 lysyl oxidase-like protein or variant thereof to be expressed in any one of the following environments including the extracellular environment, the cell surface environment, the periplasmic space or the

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intracellular environment, or a combination of any one of the environments.

- 21. A synthetic polynucleotide according to claim 20 wherein the further nucleotide sequence encodes glutathione S-transferase.
- 22. A recombinant DNA molecule comprising a synthetic polynucleotide of any one of claims 1 to 21.
- 23. A recombinant DNA molecule according to claim 22, the recombinant DNA molecule further comprising vector DNA.
 - 24. A_recombinant DNA molecule according to claim 23 wherein the vector is selected from the group consisting of pBR322, pBluescript II, pGEX-2T, pTrc99A, pET series vectors, pET3d, pCNNinv, pYEX-BX and derivatives of these vectors.
 - 25. A host cell containing a recombinant DNA molecule according to any one of claims 22, 23 or 24 or a synthetic polynucleotide according to any one of claims 1 to 21.
- 26. A host cell according to claim 25 wherein the host cell is selected from the group consisting of prokaryotic cells and eukaryotic cells.
 - 27. A host cell according to claim 26 wherein the prokaryotic cells are bacteria.
 - 28. A host cell according to claim 25 wherein the host cell is selected from the group consisting of E. coli, Saccharomyces cerevisea and Pichea.
 - 29. An expression product encoded by a synthetic polynucleotide of any one of claims 1 to 21.

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- 30. A method for producing an expression product according to claim 29, the method comprising the following steps:
- (i) providing a host cell according to any one of claims 25 to 28;

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- (ii) maintaining the host cell in conditions suitable for expression of the synthetic polynucleotide; and
 - (iii) isolating the expression product.
- 31. A method of cross-linking molecules that contain primary amines, wherein the molecules are associated with the cell membrane or the extracellular environment, the method comprising contacting the primary amine group with an expression product according to claim 29, under conditions suitable to cross-link the molecules.
 - 32. A method of cross-linking molecules that contain primary amines in vitro, the method comprising the step of contacting the primary amines with an expression product according to claim 29 under in vitro conditions which are suitable to cross-link the molecules.
 - A method of cross-linking tropoelastin, collagen or both the method comprising the step of contacting the primary amines of collagen and and/or tropoelastin with an expression product according to claim 29 under conditions which are suitable to cross-link the collagen and/or with tropoelastin.
 - A method of applying at least one cross-linkable primary amine containing molecule to a surface, wherein the surface provides primary amines available for cross-linking, the method comprising exposing the surface to the at least one primary amine containing molecule and an expression product according to claim 29 under conditions to permit the expression product to cause

cross linking of the molecules to each other and to the surface.

- 35. The method according to any one claims 31 to 34 wherein the molecule is a peptide or polypeptide and the primary amine is associated with an amino acid side chain.
- 36. The method of claim 35 wherein the peptide or polypeptide comprises the amino acid sequence of tropoelastin or a fragment thereof.
- 37. The method of claim 35 wherein the peptide or polypeptide comprises the amino acid sequence of collagen or a fragment thereof.
 - 38. The method according to any one of claims 31 to 34 wherein the molecule is a non-peptide molecule.
- 15 39. A method of accelerating the rate of wound healing, the method comprising the step of administering an expression product according to claim 29, to a wound under conditions which are suitable for cross-linking molecules which contain primary amines.
- 40. A method of accelerating the rate of wound healing according to claim 39, the method further comprising the step of administering molecules that contain primary amines to the wound under conditions which are suitable for cross-linking the molecules containing the primary amine to the wound.
 - The method according to claim 40 or 30 wherein the molecules that contain primary amines are tropoelastin molecules or derivatives thereof.

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- The method according to claim 39 or 40 wherein the molecules that contain primary amines are collagen molecules or derivatives thereof.
- 43. The method according to claim 39 or 40 wherein the molecules that contain primary amines are collagen molecules or derivatives thereof, and tropoelastin molecules or derivatives thereof.
- 44. A formulation for use in accelerating the rate of wound healing, the formulation comprising an expression product according to claim 29 and molecules containing primary amines, together with a pharmaceutically acceptable carrier, diluent or excipient.
- 45. A formulation according to claim 44 wherein the molecules containing primary amines include at least one of tropoelastin or a derivative thereof and collagen or a derivative thereof, together with a pharmaceutically acceptable carrier, diluent or excipient.
 - kit comprising a first vial containing an expression product according to claim 29 and a second vial containing molecules which contain primary amines.

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- 47. A kit according to claim 46 wherein the second vial contains tropoelastin or a derivative thereof.
- 48. A kit according to claim 46 wherein the second vial contains collagen or a derivative thereof.
- 49. A kit according to claim 47 where the kit additionally comprises a third vial containing collagen or a derivative thereof.

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- 50. A kit according to claim 46 wherein the second vial contains collagen or a derivative thereof and tropoelastin or a derivative thereof.
- 51. A method according to claim 39 or 40 wherein the expression product is administered to the wound by direct application of the product to the wound.
- 52. A method according to claim 39 or 40 wherein the expression product is administered to the wound by attaching the product to a matrix which is itself administered to the wound.
 - 53. A method according to claim 39 or 40 wherein the expression product is administered to the wound by application of a pharmaceutical formulation comprising an expression product according to claim 29 and a pharmaceutically acceptable carrier, excipient or diluent.
- 54. A method of deccelerating the rate of wound healing, the method comprising the step of administering antibodies which are raised against an expression product according to claim 29, to a wound under conditions which allow the antibodies to substantially inhibit the oxidation of primary amine moieties.
- 55. A reagent or formulation thereof for use in modifying the rate of wound healing, wherein the reagent is generated by oxidising a primary amine contained in a chemical species with an expression product according to claim 29 so as to link the chemical species to a heterologous surface.
- 56. A method of promoting fibrosis, the method comprising contacting an expression product according to claim 29 with a reagent which substantially enhances the specific acitivity of the expression product, under

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conditions which are suitable for the oxidation of primary amine moieties.

- 57. A method of promoting fibrosis, the method comprising administering an expression product according to claim 29 to a particular site under conditions suitable for the oxidation of primary amine moieties.
- 58. A method of inhibiting fibrosis, the method comprising the step of administering antibodies which are raised against an expression product according to claim 29, to a wound under conditions which allow the antibodies to substantially inhibit the oxidation of primary amine moieties.

- 59. A method for screening agents which substantially enhance or substantially inhibit the specific activity of lysyl oxidase, or lysyl oxidase-like proteins or variants thereof, the method comprising the step of contacting the agent with the expression product of the invention and measuring the specific activity of the expression product.
- 20 60. A method for detecting a protein or non-protein substrate in a sample, the method comprising contacting the sample with an expression product according to claim 29 and detecting the oxidation of any primary amine moieties in the sample.
- or non-protein substrate in a sample, wherein the number of primary amine moieties per substrate molecule is known, the method comprising contacting the sample with the expression product according to claim 29 and detecting the amount of oxidation of any primary amine moieties in the sample.

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62. A method of cancer therapy in a host in need of such treatment, comprising administering a synthetic polynucleotide according to any one of claims 1 to 21, or an expression product according to claim 29, alone or as part of a complex delivered extracellularly in the host.

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- 63. A method of regulating cell growth, the method comprising contacting a cell with an expression product according to claim 29 under conditions which permit the regulation of cell growth.
- 10 64. A method of modifying a lysine or other primary amine residue, the method comprising oxidising the primary amine by exposure to an expression product according to claim 29.
- 65. A tissue glue comprising and expression product according to claim 29 and a pharmaceutically acceptable carrier and/or diluent.
 - 66. A tissue glue according to claim 65 which additionally comprises at least one primary amine molecule species.
- 20 67. An antibody raised to an expression product according to claim 29.
 - 68. The antibody of claim 67 which is specific to an expression product according to claim 29.
- 69. A method of cross-linking molecules that contain primary amines according to claim 35, wherein the method comprises providing a complex consisting cross-linked elastin and collagen molecules, and contacting the complex with an expression product according to claim 29.
- 70. A method of accelerating the rate of wound
 30 healing, the method comprising the step of attaching the

expression product according to claim 29 to a matrix and applying the matrix to the wound.

71. The method of claim 70 wherein the matrix comprises a complex of cross-linked elastin and collagen molecules.

31/11 1/1 ATG CGC TTC GCC TGG ACC GTG CTC CTG CTC GGG CCT TTG CAG CTC TGC GCG CTA GTG CAC M R P A W T V L L G P L Q L C A L V H 91/31 TICC GOO COT COO GOO GOO GOO CAA CAG CAG COO COG CGG GAG COG GOG GOT COG GGC CAPPAAGQQPPREPPAAPG 151/51 121/41 GCC TGG CGC CAG CAG ATC CAA TGG GAG AAC AAC GGG CAG GTG TTC AGC TTG CTG AGC CTG A W R Q Q I Q W E N N G Q V F S L L S L 211/71 181/61 GGC TCA CAG TAC CAG CCT CAG CGC CGC CGG GAC CCG GGC GCC GCC CCT GGT GCA GCC G S Q Y Q P Q R R D P G A A V P G A A 271/91 241/81 AMC GOO TOO GOO CAG CAG COO COO ACT COO ATC CTG CTG ATC CGC GAC AAC CGC ACC GOO N A S A Q Q P R T P I L L I R D N R T A 331/111 GOG GOG CGA ACG CGG ACG GOC GGC TCA TCT GGA GTC ACC GCT GGC CGC CCC AGG CCC ACC TRTAGSSGVTAGRPRT AAR 391/131 361/121 GOC CGT CAC TGG TTC CAA GCT GGC TAC TCG ACA TCT AGA GCC CGC GAA CGT GGC GCC TCG T S R A R E R G A S ARHWFQAGYS 451/151 421/141 CCC GCG GAG AAC CAG ACA GCG CCC GGA GAA GTT CCT GCG CTC AGT AAC CTG CGG CCC RAENQTAPGEVPALSNL 511/171 481/161 AGC CGC GTG GAC GGC ATG GTG GGC GAC GAC CCT TAC AAC CCC TAC AAG TAC TCT GAC GAC . S R V D G M V G D D P Y N P Y K Y S D D 571/191 541/1B1 ARC COT THE THE ARC THE THE GAT HET THE GAR AGG COC AGA COT GGG GGC AGG THE COG H P Y Y H Y Y D T Y E R P R P G G R Y 631/211 CCC GGA TAC GGC ACT GGC TAC TTC CAG TAC GGT CTC CCA GAC CTG GTG GCC GAC CCC TAC G L P D L V A D P Y PGYGTGYFQY 661/221 691/231 THE ATC CAG GOG TOE ACG THE GTG CAG AAG ATG TOE ATG THE AAC CTG AGA TGC GOG GOG YIQASTYVQKHSHYNLRCA 751/251 721/241 GAG GAA AAC TGT CTG GCC AGT ACA GCA TAC AGG GCA GAT GTC AGA GAT TAT GAT CAC AGG EENCLASTAYRADVRDYDER 811/271 GTG CTG CTC AGA TIT CCC CAA AGA GTG AAA AAC CAA GGG ACA TCA GAT TIC TIA CCC AGC V L L R F P Q R V K H Q G T S D F L P S 871/291 CGA CCA AGA TAT TCC TGG GAA TGG CAC AGT TGT CAT CAA CAT TAC CAC AGT ATG GAT GAG RPRYSWEWHSCHQHYRS MDE 931/311 901/301 TITI AGC CAC TAT GAC CTG CTT GAT GCC AAC ACC CAG AGG AGA GTG GCT GAA GGC CAC AAA F S H Y D L L D A N T Q R R V A E G

961/321 991/331 GCA AGT TTC TGT CTT GAA GAC ACA TCC TGT GAC TAT GGC TAC CAC AGG CGA TTT GCA TGT A S F C L E D T S C D Y G Y H R R F A C 1021/341 1051/351 ACT GCA CAC ACA CAG GGA TTG AGT CCT GGC TGT TAT GAT ACC TAT GGT GCA GAC ATA GAC T A H T Q G L 6 P G CYDTYGADID 1081/361 1111/371 TGC CAG TGG ATT GAT ATT ACA GAT GTA AAA CCT GGA AAC TAT ATC CTA AAG GTC AGT GTA C Q W I D I T D V K P G N Y I L K V S V 1141/381 1171/391 AAC CCC AGC TAC CTG GTT CCT GAA TCT GAC TAT ACC AAC AAT GTT GTG CGC TGT GAC ATT N P S Y L V P E S D Y T N N V V R C D I 1201/401 1231/411 CGC TAC ACA GGA CAT CAT GCG TAT GCC TCA GGC TGC ACA ATT TCA CCG TAT R Y T G H H A Y A S G C T I S P Y

Fig. 1(2)

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GA TCC CAG CAG CAG CCG CCG CGT GAA CCG CCG GCT GCT CCG GGT GCT TGG CGT CAG CAG

S Q Q Q P P R E P P A A P G A W R Q Q

61/21

ATC CAG TGG GAA AAC AAC GGT CAG GTT TTC TCC CTG CTG TCC CTG GGT TCC CAG TAC CAG

I Q W E N N G Q V F E L L E L G E Q Y Q

121/41

CCG CAG CGT CGT CGT GAC CCG GGT GCT GCT GTT CCG GGT GCT GCT CAG GCT TCC GCT CAG

P Q R R D P G A A V P G A A N A 6 A Q

181/61

211/71

CAG CCG CGT ACC CCG ATC CTG CTG ATC CGT GAC AAC CGT ACC GCG GCC GCT CGT ACC CGT Q P R T P I L L I R D N R T A A A A R T R

PstI

241/81

271/91

ACC GCT GGT TCC TCC GGT GTT ACT GCA GGT CGT CCG CGT CCG ACC GCG CGC CAC TGG TTC

T A G E E G V T A G R P R P T A R R H W F

301/101

CAG GCT GGT TAC TCC ACC TCC CGT GCT CGT GAA GCT GGT GCT TCC CGT GAA AAC CAG

Q A G Y E T E R A R E A G A E R A E N Q

361/121

ACC GCT CCG GGT GAA GTT CCA GCG CTG TCC AAC CTG CGT CCG CGT CCG CGT CCG CGT GCT GAC GGT

T A P G E V P A L E N L R P P E R V D G

ATG OTT GGT GAC GAC COG TAC AAC COG TAC AAG TAC TOC GAC GAC AAC CCG TAC TAC AAC

KpnI

481/161

421/141

511/171

451/151

TAC TAC GAC ACC TAC GAG CGC CGG CGT CCG GGT GGT CGT TAC CGT CCG GGT TAC GGT ACC

Y Y D T Y E R P R P G G R Y R P G T

541/181

GGT TAC TTC CAG TAC GGT CTG CCG GAC CTG GTT GCT GAC CCG TAC TAC ATC CAG GCT TCC
G Y F Q Y G L P D L V A D P Y Y I Q A S

Fig. 2(1)

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751/251

CCG CAG CGT GTT AAA AAC CAG GGC ACC TCC GAC TTC CTG CCG TCC CGT CCG CGT TAC TCC
P Q R V K N Q G T S D F L P S R P R Y S

Ecori

781/261

721/241

811/271

TGG GAA TGG CAC TCC TGC CAC CAG CAC TAC CAC TCC ATG GAC GAA TTC TCC CAC TAC GAC

W E W E S C E Q E Y E S M D E F 6 H Y D

841/281

CTG CTG GAC GCT AAC ACC CAG CGT CGT GTT GCT GAA GGT CAC AAA GCT TCC TTC TGC CTG

L L D A N T Q R R V A E G H K A E F C L

901/301

GAA GAC ACC TCC TGC GAC TAC GGT TAC CAC CGT CGT TTC GCT TGC ACC GCT CAC ACC CAG \mathbb{R}^2 $\mathbb{$

GGT CTG TCC CCG GGT TGC TAC GAC ACC TAC GGT GCT GAC ATC GAC TGC CAG TGG ATC GAC

G L S P G C Y D T Y G A D I D C Q W I D

1021/341

ATC ACC GAC GIT AAA CCG GGT AAC TAC ATC CTG AAA GIT TCC GIT AAC CCG TCC TAC CTG

I T D V K P G N Y I L K V 6 V N P 6 Y L

1081/361

GTT COG GAA TCC GAC TAC ACC AAC AAC GTG GTT CGT TGC GAT ATC CGT TAC ACC GGT CAC

V P E S D Y T N N V V R C D I R Y T G H

BamHI

1141/381

1171/391

CAC GCT TAC GCT TCC GGT TGC ACC ATC TCC CCG TAC TAA TGA TAG TAG H A Y A S G C T I S P Y * * *

Fig. 2(2)

81 CCAACAGCAGCCCCGCGCGAGCCGCCGGCGGCGCGCCCCGGGCGCCCCCC
6 CCAGCAGCAGCCGCCGCGTGAACCGCCGGCTGCTCCGGGTGCTTGGCGTC 55
131 AGCAGATCCAATGGGAGAACAACGGGCAGGTGTTCAGCTTGCTGAGCCTG 180
56 AGCAGATCCAGTGGGAAAACAACGGTCAGGTTTTCTCCCTGCTGTCCCTG 105
181 GGCTCACAGTACCAGCCTCAGCGCCCGGGGACCCGGGCGCCGCCGTCCC 230
106 GGTTCCCAGTACCAGCCGCAGCGTCGTCGTGACCCGGGTGCTGCTCC 155
231 TGGTGCAGCCAACGCCTCCGCCCAGCAGCCCCGCACTCCGATCCTGCA 280
281 TCCGCGACAACCGCACCGCCGCGCGCGCGACGGCCGGCTCATCT 330
331 GGAGTCACCGCTGGCCGCCCCAGGCCCACCGCCCGTCACTGGTTCCAAGC 380
381 TGGCTACTCGACATCTAGAGCCCGCGAACGTGGCGCCTCGCGCGCG
431 ACCAGACAGCGCCGGGAGAAGTTCCTGCGCTCAGTAACCTGCGGCCGCCC 480
481 AGCCGCGTGGACGCCATGGTGGGCGACGACCCTTACAACCCCTACAAGTA 530
531 CTCTGACGACAACCCTTATTACAACTACTACGATACTTATGAAAGGCCCA 580
581 GACCTGGGGGCAGGTACCGGCCCGGATACGGCACTGGCTACTTCCAGTAC 63
631 GGTCTCCCAGACCTGGTGGCCGACCCCTACTACATCCAGGCGTCCACGTA 68



WO 98/06830

681	CGTGCAGAAGATGTCCATGTACAACCTGAGATGCGCGGGGGGAAAACT	730
606	CGTTCAGAAAATGTCCATGTACAACCTGCGTTGCGCTGCTGAAGAAACT	655
731	GTCTGGCCAGTACAGCATACAGGGCAGATGTCAGAGATTATGATCACAGG	780
656		705
781	GTGCTGCTCAGATTTCCCCAAAGAGTGAAAAACCAAGGGACATCAGATTT	830
706		755
831	CTTACCCAGCCGACCAAGATATTCCTGGGAATGGCACAGTTGTCATCAAC	880
756	CCTGCCGTCCGCGTTACTCCTGGGAATGGCACTCCTGCCACCAGC	805
881	ATTACCACAGTATGGATGAGTTTAGCCACTATGACCTGCTTGATGCCAAC	930
806	ACTACCACTCCATGGACGAATTCTCCCACTACGACCTGCTGGACGCTAAC	855
931	ACCCAGAGGAGAGTGGCTGAAGGCCACAAAGCAAGTTTCTGTCTTGAAGA	980
856	ACCCAGCGTCGTGTTGCTGAAGGTCACAAAGCTTCCTTCTGCCTGGAAGA	905
981	CACATCCTGTGACTATGGCTACCACAGGCGATTTGCATGTACTGCACACA	1030
906	CACCTCCTGCGACTACGGTTACCACCGTCGTTTCGCTTGCACCGCTCACA	955
1031	CACAGGGATTGAGTCCTGGCTGTTATGATACCTATGGTGCAGACATAGAC	1080
	CCCAGGGTCTGTCCCCGGGTTGCTACGACACCTACGGTGCTGACATCGAC	1005
1081	TGCCAGTGGATTGATATTACAGATGTAAAACCTGGAAACTATATCCTAAA	1130
	TGCCAGTGGATCGACATCACCGACGTTAAACCGGGTAACTACATCCTGAA	
	GGTCAGTGTAAACCCCAGCTACCTGGTTCCTGAATCTGACTATACCAACA	
	AGTITCCGTTAACCCGTCCTACCTGGTTCCGGAATCCGACTACACCAACA	
	ATGTTGTGCGCTGTGACATTCGCTACACAGGACATCATGCGTATGCCTCA	
	ACGTTGTTCGTTGCGATATCCGTTACACCGGTCACCACGCTTACGCTTCC	1155
	GCTGCACAATTTCACCGTAT 1251	
	~ ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	

Fig. 3(2)

101/10///000

BLOCK 1: Coding strand

Oligo 1

GATCCCAGCAGCAGCCGCCGCGTGAACCGCCGGCTGCTCCGGGTGCTTGGCGTCAGCAGATCCAGTGGGAAAACAACGGTCAGGTTTTCTCCCTG (SEQ ID NO: 5)

Oligo 2

pcTGTCCCTGGGTTCCCAGTACCAGCCGCAGCGTCGTGACCCGGGTGCTGCTGCTCCGGGTGCTGCTAACGCTTCCGCTCAGC (SEQ ID NO: 6)

Oligo 3

PAGCCGCGTACCCCGATCCTGCTGATCCGTGACAACCGTACCGCGCCGCTCGTACCCGTACCCGTACCCGTGTTACTGCA (SEQ ID NO: 7)

BLOCK 2: Coding strand

Oligo 4

GGTCGTCCGCGCCGCGCGCCACTGGTTCCAGGCTGGTTACTCCACCTCCCGTGCTCGTGAAGCTGGTGCTTCCCGTGCTGAAAACCAG (SEQ ID NO: 8)

Oligo 5

PACCGCTCCGGGTGAAGTTCCAGCGCTGTCCAACCTGCGTCCGCCGTCCCGTGTTGACGGTATGGTTGGTGACGACCCGTACAACCCGTACA (SEQ ID NO: 9)

Oligo 6

PAGTACTCCGACGACACCCGTACTACAACTACTACGACACCTACGAGCGCCCGCGTCCGGGTGGTCGTTACCGTCCGGGTTACGGTAC (SEQ ID NO: 10)

BLOCK 3: Coding strand

Oligo 7

CGGTTACTTCCAGTACGGTCTGCCGGACCTGGTTGCTGACCCGTACTACATCCAGGCTTCCACCTACGTTCAGAAAATGTCCATGTAC (SEQ ID NO: 11)

Oligo 8

PAACCTGCGTTGCGCTGAAGAAACTGCCTGGCTTCCACCGCTTACCGTGCTGACGTTCGTGACTACGACCACCGTGTTCTGCTGCGTTTCCCGC (SEQ ID NO: 12)

Fig. 4(1)

8/12

Oligo 9

11 C 20/00030

PAGCGTGTTAAAAACCAGGGCACCTCCGACTTCCTGCCGTCCCGCGTTACTCCTGG-GAATGGCACTCCTGCCACCAGCACTACCACTCCATGGACG (SEQ ID NO: 13)

BLOCK 4: Coding strand

<u>Oligo</u> 10

AATTCTCCCACTACGACCTGGACGCTAACACCCAGCGTCGTGTTGCTGAAGGTCACA-AAGCTTCCTTCTGCCTGGAAGACACCTCCTGCGACTACG (SEQ ID NO: 14)

<u>01igo 11</u>

PGTTACCACCGTCGTTTCGCTTGCACCGCTCACACCCAGGGTCTGTCCCCGGGTTGCTAC-GACACCTACGGTGCTGACATCGACTGCCAGTGGA (SEQ ID NO: 15)

<u>Oligo 12</u>

PTCGACATCACCGACGTTAAACCGGGTAACTACATCCTGAAAGTTTCCGTTAACCCGTCC-TACCTGGTTCCGGAATCCGACTACACCAACAACG (SEQ ID NO: 16)

Oligo 13

PTGGTTCGTTGCGATATCCGTTACACCGGTCACCACGCTTACGCTTCCGGTTGCACCATC-TCCCCGTACTAATGATAG (SEQ ID NO: 17)

COMPLEMENTARY OLIGONUCLEOTIDES

BLOCK 1: Complementary strand

Complementary to oligo 1

PGAAAACCTGACCGTTGTTTTCCCACTGGATCTGCTGACGCCAAGCACCCGGAGCAGCCG-GCGGTTCACGCGGCGGCTGCTGG (SEQ ID NO: 18)

Complementary to oligo 2

PCGGAAGCGTTAGCAGCACCCGGGAACAGCACCCGGGTCACGACGACGCTGCGGCTGG-TACTGGGAACCCAGGGACAGCAGGGA (SEQ ID NO: 19)

Fig. 4(2)

Complementary to oligo 3

GTAACACCGGAGGAACCAGCGGTACGGGTACGGGTACGGGTACGGGTTGTCACGGATC-AGCAGGATCGGGGTACGCGGCTGCTGAG (SEQ ID NO: 20)

BLOCK 2: Complementary strand

Complementary to oligo 4

PTTCAGCACGGGAAGCACCAGCTTCACGAGCACGGGAGGTGGAGTAACCAGCCTGGAACCAGTGGCGCGCGGTCGGACGCGGACGACCTGCA (SEQ ID NO: 21)

Complementary to oligo 5

PGTTGTACGGGTCGTCACCAACCATACCGTCAACACGGGACGGGGGGGCGGACGCAGGTTGGACA-GCGCTGGAACTTCACCCGGAGCGGTCTGGTT (SEQ ID NO: 22)

Complementary to oligo 6

CGTAACCCGGACGGTAACGACCACCCGGACGCGGGCGCTCGTAGGTGTCGTAGTTGTAGTACGGGTTGTCGTCGGAGTACTTGTACGG (SEQ ID NO: 23)

BLOCK 3: Complementary strand

Complementary to oligo 7

pgcaggttgtacatggacattttctgaacgtaggtggaagcctggatgtagtacgggtcagcaaccaggtccggcagaccgtactggaagtaccggtac (seq id no: 24)

Complementary to oligo 8

PACACGCTGCGGAAACGCAGCAGAACACGGTGGTCGTAGTCACGAACGTCAGCACGGTAAGCGGTGGAAGCCAGGCAGTTTTCTTCAGCAGCGCAAC (SEQ ID NO: 25)

Complementary to oligo 9

AATTCGTCCATGGAGTGGTAGTGCTGGTGGCAGGAGTGCCATTCCCAGGAGTAACGCGGA-CGGGACGCAGGAAGTCGGAGGTGCCCTGGTTTTTA (SEQ ID NO: 26)

Fig. 4(3)

BLOCK 4: Complementary strand

Complementary to oligo 10

PGCAGGAGGTGTCTTCCAGGCAGAAGGAAGCTTTGTGACCTTCAGCAACACGACGCTGGGTGTTAGCGTCCAGCAGGTCGTAGTGGGAG (SEQ ID NO: 27)

Complementary to oligo 11

pGGCAGTCGATGTCAGCACCGTAGGTGTCGTAGCAACCCGGGGACAGCCCTGGGTGTGA-GCGGTGCAAGCGAAACGACGGTGGTAACCGTAGTC (SEQ ID NO: 28)

Complementary to oligo 12

PTGGTGTAGTCGGAACCAGGTAGGACGGGTTAACGGAAACTTTCAGGATGTAGTTACCCGGTTTAACGTCGGTGATGTCGATCCACT (SEQ ID NO: 29)

-Complementary to oligo 13

GATCCTATCATTAGTACGGGGAGATGGTGCAACCGGAAGCGTAAGCGTGGTGACCGGTGTAACGGATATCGCAACGAACCACGTTGT (SEQ ID NO: 30)

Fig. 4(4)

11/12

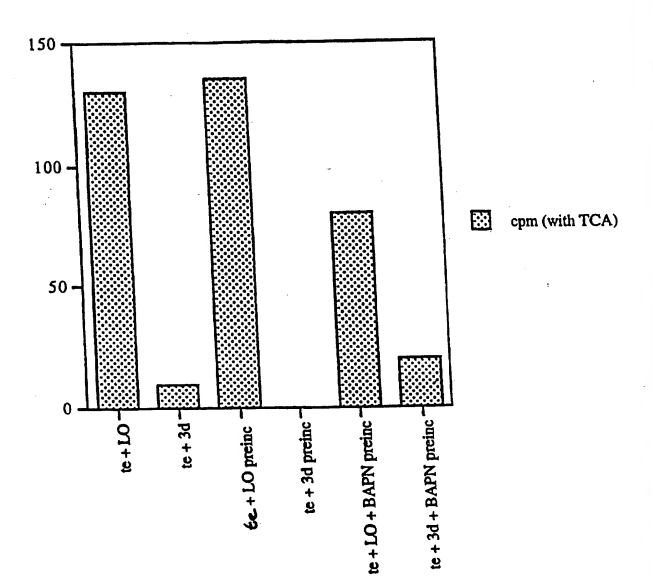


Fig. 5

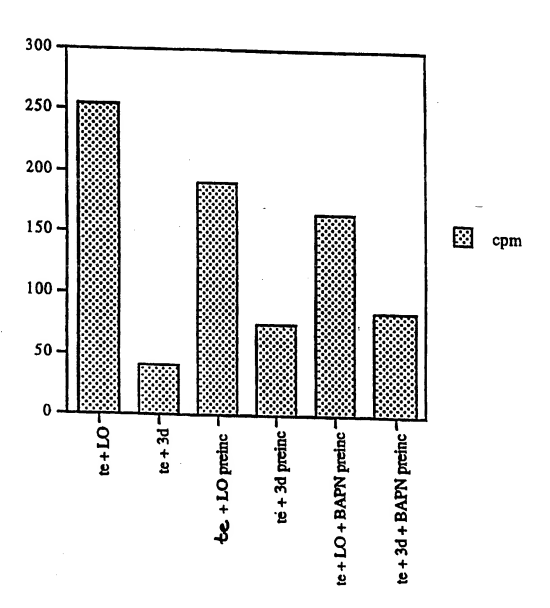


Fig. 6

Authorized officer

MADHU K. JOGIA

Telephone No.: (02) 6283 2512

mjogia

Name and mailing address of the ISA/AU

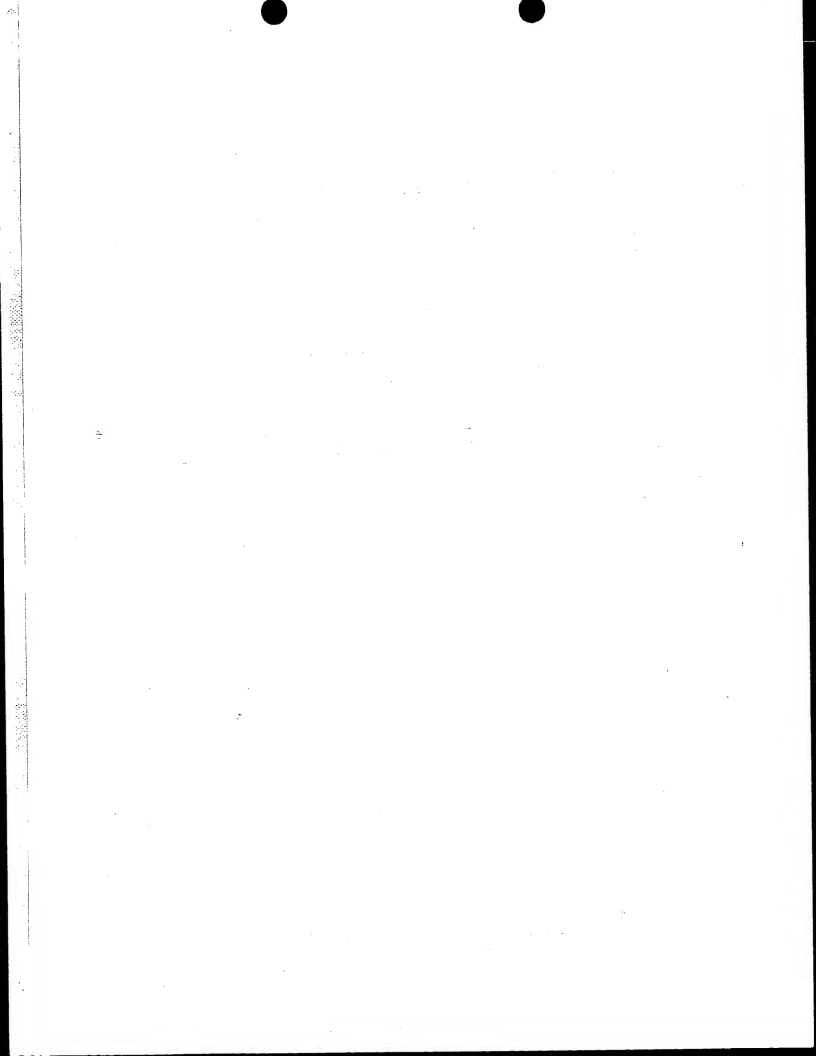
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Facsimule No.: (02) 6285 3929

C (Continua	PCT/AU 97/00505	
	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	J. Biol. Chem, vol. 268(25), pp 18435-18437, published sept. 5, 1993. Kenyon, k et al. "A novel human CDNA with a predicted protein similar to lysyl oxidase maps to chromosome 15q24-q25".	1, 2, 3, 17, 18 19-23.
X	Genomics, vol. 17(3), pp 544-548, published sept. 1993. Hamalainen et al. "structure of the human lysyl oxidase gene".	1, 2, 3, 17, 1 19-23.
X	J. Biol. Chem, vol. 270(13), pp 7176-7182, published March 31, 1995.kim, Y.et al. "A new gene with sequence and structure similarity to the gene encoding human lysyl oxidase'.	1, 2, 3, 17, 1 19-23
P,X	FEBS Lett, vol 399(3), pp 215-219 published Dec 16, 1996. Ouzzine, M. et al. "Expression of active, human lysyl oxidase in Escherichia coli".	1, 2, 3, 5
X	Matrix, vol.12, pp 242-248, published 1992 Mariani, T.J.et al. "The complete derived amine acid sequence of human lysyl oxidase and assignment of the gene to chromosome 5"	
X	JP, 04-021700 (NIPPON KAYAKU) 24 Jan, 1992.	1, 2, 3, 5, 17, 18, 19, 2
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1/1 31/11 ANG COST THE GOOD TOOK ACCOUNTS CITE CITE CITE GOOD COT TITE CAME CITE THE GOOD CITA GITE CIAC ERFANTVLLLGPLOLOALVE 61/21 91/21 THE GOO DOT COO GOO GOO GOO CHA CAL CAL CAL COC COO COO CAL CAL COO COO GOO GOT COO CAL 121/41 151/51 GOO TIGG CGC CAG CAG ATC CAA TOG GAG AAC AAC GGG CAG GTG TTC AGC TTG CTG AGC CTG A W R Q Q I Q W E R N O Q V F E L L E L 211/71 GRO TON ONE THE CAS COT CAS GRO COS CAS COS GRO COT GRO GRO COT GRT GRA GRO G 6 Q Y Q P O R R D P G A A V P G A A 241/81 271/91 ANC GOO TOO GOO CAG CRAS COO COO ACT COO ATC CTG CTG ATC CGC GAC AND CGC ACC GOO R A G Q P R T P ILLIRD 301/101 331/111 DOG GOG COR ACC CGG ACG GOC GOC TOR TOT GGR GTC ACC GOT GGC CGC CCC AGG CCC ACC AARTRTAGESGVTAGRPRP 361/121 391/131 DOG COT CAC TOG THE CAR GET COC THE TOT ACA TOT AGA GEO COC GAA COT GGC GCC_TOG ARRWFQAGYS-TSRARRGAS 421/141 491/151 COC GOO GAG AAC CAN ACA GOO COC GOA GAA GIT COT GOO CIC AGI AAC CIG COO COC COC RAERQTAFGEVPALSULRPP 481/161 511/171 AGO OGC GRIC GAC GGC ATG GRIC GGC GAC GAC CCT TAC AAC CCC TAC AAG TAC TCT GAC GAC . еклодидертивькиер р 541/101 571/191 ARC COT THE THE ARC THE SHE GRE NOT THE GRA HOS COO AGE COT GGG GGC AGG THE COG R P Y T H Y Y D T Y E R P R P G G R Y R 601/201 631/211 CCC GRA TRO GGC ACT GGC TRO TRO CMG TRO GGT CTC CCA-GRA CTG GTG GCC GRA CCC TRO PGYGTGYFOLVAD 661/221 691/231 THE ARE CAG GOS TOO ACS THE STS CAS ARE ATS TOO ATS THE THE ARE CTG AGA TOO GOS GOS Y I Q A S T Y Y Q K N S H T H L R C A A 731/251 GAG GAA AAC TOT CITY GCC ANT ACA GCA TAC AGG GCA GRIT GTC AGA GRIT TAT GAT CAC AGG B C B C L A S T A T R A D V R D Y D E 781/261 811/271 GRE CHE CHE AGA THE COC CAR AGA CHE AAR AMC CAR GOG AGA TER GAT THE THR COC AGE V L L R P P Q R V K B Q G T B D F L P S 871/291 COR CUL MOR TRI TOC TOO GRA TOG CAC AGT TOT CAT CAR CRI TAC CAC AGT ATG GAT GAG RPRYSUSHBCEQUYESHDE 931/311 THE AGO CAC THE GAC CHE CHE GAT GOD AND ACC CAG AGG AGA GHO GOT GAA GGC CAC AAA PEGYDLLDANTORRVAEGEK

991/331 961/321 SCA ACT THE TOT CIT GAA GAE ACA TOE TOT GAE TAT GGC TAE CAE AGG CGA THT GCA TOT ASPCLEDTSCDYGTHARPAC 1051/351 1021/341 ACT GCA CAC ACA CAG GGA TTG AGT CCT GGC TGT TAT GAT ACC TAT GGT GCA GAC ATA GAC TARTQGLEPGCYDTTGADI 1111/371 1081/361 TOC CAS TOG ATT GAT ATT ACA GAT GTA ARA CCT GOA ARC TAT ATC CTA ARG GTC AGT GTA C Q W I D I T D V K P G H Y I L K V S V 1171/391 1141/381 AME OCC AGE THE CTG GTT CET GAR TET GRE THT MOE ARE BAT GTT GTG OGC TOT GAE ATT R P S Y L Y P E S D Y T R R Y Y R C D I 1231/411 1201/401 CON TAC ACA GGA CAT CAT GCG TAT GCC TCA GGC TGC ACA ATT TUA DCG TAT RYTGHEAXAGGCTISFY

Fig. 1(2)

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2/

31/11

ATC CAG TOG GAN AND AND GOT CAG GTT TTC TOC CTG GTG TOC CTG GOT TOD CAG TAG CAG . . .

I Q W E W F G Q V F B L L G L G S Q Y Q 121/61

CAS COG COT ACC COG ATC CTG CTG ATC COT GAC AAC CGT ACC GCT CCT CCT ACC CGT
Q P R T P I L L I R D N R T A A A R T R

PatI

241/81

271/91

ACC GCT GGT TCC GGT GTT ACT GCA GGT CCT CCG GCT CCG ACC GCG CGC CAC TGG TTC

T A G E E G V T A G R P R P T A R E W F

301/101

CAG GCT GGT TAG TOG ACC TOC CGT GCT GCT GGT GGT GGT GCT TOC CGT GCT GAA AAC CAG
Q A G Y S T S R A R E A G A S R A E R Q
361/121

ACC OCT COG GOT GAA GTT CCA GCG CTG TCC AAC CTG CGT CCG CCG TCC CGT GTT GAC GGT

T A P G E V P A L 8 N L R P P 8 R V D G

451/151

ATG GET GAT GAT GAT GOG TAC AAC CCG TAC AAG TAC TCC GAT GAT AAC CCG TAC TAC AAC

Exal

681/161

511/171

THE TAC GARD ALSO TIME GAME COSC COST COST COST GIGT COST TRAC COS

GOT THE THE CAS THE GOT CHG COC GAS CHG OTT GOT GAS COG THE THE ATC CAS GOT THE G Y F Q Y G L P D L V A D P Y Y I Q A S

Fig. 2(1)

501/201

631/211

ACC TAC STT CAG AAA ATG TCC MTG TAC AAC CTG CST TGC GCT GCT GAA GAA AAC TGC CTG

T Y V Q K M E H Y H L R C A A E E H C L

661/221

691/231

COL CAG COT GIT MAR MAC CAG GGC ACC TOC GAC TIC CTG COO TOC CGT CCG CON TAC TOC P Q R V K N Q G T S D F L P B R P R Y S

EcoRI

781/261

811/271

TOG GAA TOG CAC TOC TOC CAC CAC CAC CAC CAC TOC ATC GAC GAA TTC TOC CAC TAC GAC

W E W E S C B Q E Y E S N D E F S B Y D

861/281

CING CING GROUP SAME ARCHITECT CAST COTT GROUP CARC GROUP CARC ANA GROUP CARC AND GROUP CARC CINC AND GROUP CARC AND GROUP CARC CINC AND GROUP CARC AND GROU

CAN CAR ACC TOC TOC CAR TAR COT THE CAR COT TOT COT TOT ACC COT CAR ACC CAR

B D T S C D Y G Y B R R F A C T A B T Q

961/321

991/331

001 CTG TCC CC3 G32 TGC TAC GAC ACC TAC G37 GCT GAC ATC GAC TGC CAG TGG ATC GAC
G L 8 P G C Y D T Y G A D I D C Q W I D
1021/341

AND AND GAR STT ARA COO GET AND THE MIC ONG ARA STT TOO OUT AND COS TOO THE CITY I \hat{T}^{ab} by K p g H y I L K V B V N P S I L 1081/361

OFF COO GAR TOO GAR! TAC ACC AAC AAC ONG GIT COT TOO GAR AND CON TAC ACC GOT CAC

BennHI

1141/381

1171/391

CAC GCT TAG GCT TOC GGT THE ACC ATC TOC CCG TAG TAA THA THA TAG B A Y A S G C T I B P Y + * *

Fig. 2(2)

	CCALCAGERGOCCCGCGCGGASCCGCCGGGCGGCGCGCGCGCGCGCGCGCG	
	CCAGCAGCAGCCGCGCGGAACCGCCGGGTGCTTGGCGTC	
	AGCAGATCCAATGGGAGAACAACGGGCAGGTGTTCAGCTTGCTGAGCCTG	
	AGCAGATCCAGTGGGAAACAACGGTCAGGTTTTCTCCCTGCTGTCCCTG	
	GGCTCACAGTACCAGCCTCAGCGCCGCGGGGGGGGGGGG	
	CGTTCCCAGTACCAGCCCCAGCGTCGTCGTCACCCGGGTGCTGCTCTCC	
	TGGTGCAGCCAACGCCTGCGCCCAGCAGCCGCGCACTCCGATCCTGCTGA	
	GGGTGCTGACGCTTCCGCTCAGCAGCCGCGTACCCCGATCCTGCTGA	
281	TCCGCGACACCGCACCGCGCGCCGCACGCCGCCCCCCCCC	330
	TCCGTGACAACCGTACCGCGCCCCCTCGTACCGCTGCTTCCTCC	
331	GGAGTCACCCCGGCCGCCCCAGGCCCACCGGCCCCTCACTGGTTOCAAGC	380
256		305
301	TGGCTACTCGACATCTAGAGCCCGCGCAACGTGGCGCGCGC	430
306	TEGTTACTCCACCTCCCGTGCTCGTGAAGCTGCTGCTCCCGTGCTGAAA	355
431	ACCAGACAGEGGGGGGGGAGAACTTCCTGCGCTCAGTAACCTGCGGCCGCCCC	480
356	ACCAGACCGCTCCGGGTGAAGTTCCAGCGCTGCCAACCTGCGTCCGCCG	405
481	AGCCGCGTGGACGGCATGGTGGGGGGGGGGGCCCTACAAGTA	530
406	TOCCGTGTTGACGGTATGGTTGGTGACGACCGGTACAACTA	455
531	CTCTGACCACCACCCTTATTACAACTACTACGATACTTATGAAAGGCCCA	5B0
456	CTCCGACGACAACCCGTACTACAACTACGACACCTACGAGCGCCCGC	505
581	GACCTGGGGGCAGGTACCGGCCCGGCTACTTCCAGTAC	630
	CACCAGOTEGTCACCGTCCGGGGTTACGGTTACGTTACTTCCAGTAC	555
	GGTCTCCCAGACCTGGTGGCCGACCCCTACTACATCCAGGCGTCCACGTA	
556	GGTCTGCCGGACCTGGTTGCTGCCCGTACTACATCCAGGCTTCCACCTA	604

81 CGTGCAGAAGATGTCCATGTACAACCTGAGATGCGCGGGGGGGG
731 GYCTGGCCAGTACAGCATACAGGGCAGATGTCAGAGATTATGATCACAGG 780
656 GCCTGGCTTCCACCGCTTACCGTGCTGACGTTCGTGACTACGACCACCGT 705
781 GTGCTGCTCAGATTTCCCCCAAAQAGTGAAAACCAAGGGACATCAGATTT 830
706 CITCTGCTGCGTTTCCCGCAGCGTGTTAAAAACCAGGGTACCTCCGACTT 155
831 CTTACCOACCCGACCAAGATATTCCTGGGALTGGCACAGTTGTCATCAAC 860
756 CCTGCCGTCCCGTCGCGTTACTCCTGGGAATGGCACTCCTGCCACCAGC 805
BEI ATTACCACAGTATGGATGAGTTTAGCCACTATGACCTGCTTGATGCCAAC 930
#
931 ACCCAGAGGAGAGTGGCTGAAGGCAAGGAAGTTTCTGTCTTGAAGA 980
B56 ACCCAGCGTCGTGCTGAAGGTCACAAAGCTTCCTTCTGCCTGGAAGA 905
981 CACATCOTGREACTATGGCTACCACAGGCGATTTGCATGTACTGCACACA 1030
906 CACCTCCTGCGACTACGGTTACCACCGTCGTTTCGCTTGCACCGCTCACA 955
1031 CACAGGGATTGAGTCCTGGCTGTTATGATACCTATGGTGCAGACATAGAC 1080
956 CCCAGGGTCTGTCCCCGGGTTGCTACGACACCTACGGTGCTGACATCGAC 1095
1081 TGCCAGTGGATTGATATTACAGATGTANAACCTGGAAACTATATCCTAAA 1130
1006 TGCCAGTGATCGACATCACCGACGTTAAACCGGGGTAACTACATCCTGAA 1055
1131 GGTCAGTGTAAACCCCAGCTACCTGGTTCCTGAATCTGACTATACCAAGA 1180
1056 AGTTTCCGTTAACCCGTCCTACCTGGTTCCGGAATCCGACTACACCAACA 1105
1181 ATGITGTGCGCTGTGACATTCGCTACACGGACATCATGCGTATGCCTCA 1231
4
1231 GGCTGCACAATTTCACCGTAT 1251
1152 <i>ማለዘነስሮሽኒ ሲ</i> ለይመለክቸውም/የመከት 1176

7,

BLOCK 1: Coding strand

Oligo 1

GATCCCAGCAGCAGCAGCGCGGGTGAACCGCCGGCTGCTCCGGGTGCTTGGCGTCACCAGA—
TCCAGTGGGAAAACAACGGTCAGGTTTTCTCCCCTG (SEQ ID NO: 5)

Oligo 2

Oligo 3

PAGCCGCGTACCCGATCCTGATCCGTGACAACCGTACCGCGCCCCCCGTACCCGT-ACCGCTGCTACCCGTGTTACTGCA (SEQ ID NO: 7)

BLOCK 2: Coding strand

<u>011go 4</u>

GGTCGTCCGCGTCCGACCGCGCCACTGGTTCCAGGCTGGTTACTCCACCTCCCGTGCTCGTGAAGCTGGTGCTTCCCGTGCTGAAAACCAG (SEQ ID NO: B)

Olico 5

PACCGCTCCGGGTGAAGTTCCAGCGCTGTCCAACCTGCGTCCGCCGTGTTGACGGTATGGTTGGTGACGACCCGTACAACCCGTACA (SEQ ID NO: 9)

<u>Oliao 6</u>

PAGTACTCCGACGACACCCGTACTACGACTACGACACCTACGAGCGCCCGCGTCCGGGTGGTCGTTACCGTCCGGGTTACGGTAC
(SEQ ID NO: 10)

BLOCK 3: Coding strand

<u> Oligo 7</u>

CGCTACTTCEAGTACGGTCTGCCGGACCTGGTTGCTGACCCGTACTACATCCAGGCTTC-CACCTACGTTCAGAAAATGTCCATGTAC (SEQ ID NO: 11)

Olico 8

PAACCTGCGTTGCGCTGAGAAAACTGCCTGGCTTCCACCGCTTACCGTGCTGACGTTCGTGACTACGACCACCGTGTTCTGCTGCGTTTCCCGC (SEQ ID NO: 12)

Fig. 4(1)

Oligo 9

BLOCK 4: Coding strand

<u> 01ias 10</u>

AATTCTCCCACTACGACCTGGACGCTAACACCCCAGCGTCGTGTTGCTGAAGGTCACAAAGCTTCCTTCTGCCTGGAAGACACCTCCTGCGACTACG (SEQ ID NO: 14)

01igo 11

pGTTACCACCGTCGTTTCGCTTGCACCCCCACACCCAGGTCTGTCCCCGGGTTGCTAC-GACACCTACGGTCCTGACATCGACTGCCAGTGGA (SEQ ID NO: 15)

011Go 12

pTCGACATCACCGACGTAACCGGGTAACTACATCCTGAAAGTTTCCGTTAACCCGTCC-TACCTGGTTCCGGAATCCGACTACACCAACACG (SEQ ID NO: 16)

011co 13

prestrected technique (SEQ ID NO: 17)

COMPLEMENTARY OLIGONUCLEOTIDES

BLOCK 1: Complementary strand

Complementary to oligo 1

Complementary to oligo 2

Fig. 4(2)

Complementary to olico 3

GTAACACCGGAGGAACCAGCGGTACGGGTACGGGTACGGGTACGGGATC-AGCAGGATCGGGGTACGCGGCTGCTGAG (SEQ ID NO: 20)

BLOCK 2: Complementary strand

Complementary to oligo 4

PTTCAGCACGGGAGCACCAGCTTCACGAGCACGGGAGGTGGAGTAACCAGCCTGGAACC-AGTGGCGCGCGGGCGGACGGGACGTGCA (SEQ 1D NO: 21)

Complementary to oligo 5

Complementary to oligo 6

BLOCK 3: Complementary atrand

Complementary to plice 7

PGCAGGTTGTACATGGACATTTTCTGAACGTAGGTGGAAGCCTGGATGTACGGGTCA-GCAACCAGGTCCGGCAGACCGTACTGGAAGTACCGGTAC (SEQ 1D NO: 24)

Complementary to plice 8

Complementary to oligo 9

AATTCGTCCATGCAGGAGTGCTGGTGGCAGGAGTGCCATTCCCAGGAGTAACGCGGA-CGGGACGCAGGAAGTCGGAGGTGCCCTGGTTTTTA (SEQ ID NO: 26)

Fig. 4(3)

BLOCK 4: Complementary strand

Complementary to oligo 10

pgcaggaggtgtcttccaggcagaaggaagctttgtgaccttcagcaacacgacgctgggtgttagcgtccagcaggtcgtagtgggag (SEQ ID NO: 27)

Complementary to oligo 11

pggcagtcgatgtcagcaccgtaggtgtcgtagcaacccggggaCagaccctgggtgtgagcggtgcaagcgaaacgacggtggtaaccgtagtc (SEQ ID NO: 28)

Complementary to oligo 12

pTGGTGTAGTCGGATCCGGAACCAGGTAGGACGGTTAACGGAAACTTTCAGGATGTAGTTACCCGGTTTAACGTCGGTGATGTCGATCCACT (SEQ ID NO: 29)

Complementary to oligo 13

GATCCTATCATTAGTACGGGGAGATGGTGCAACCGGAAGCGTAAGCGTGACCGGTGT-AACGGATATCGCAACGAACCACGTTGT (SEO ID NO: 30)

Fig. 4(4)

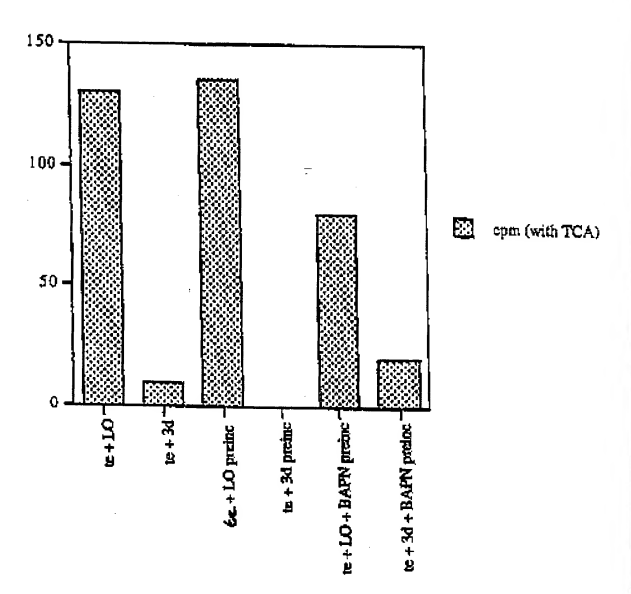


Fig. 5

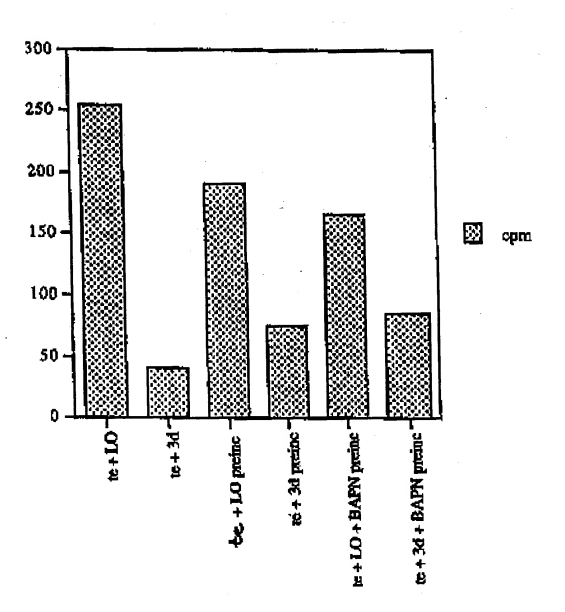


Fig. 6